



Impact of impurities on water electrolysis: a review

Cite this: *Sustainable Energy Fuels*,
2023, 7, 1565

Hans Becker,^a James Murawski,^b Dipak V. Shinde,^a Ifan E. L. Stephens,^{b*}
Gareth Hinds^a and Graham Smith^{a*}

Low temperature water electrolyzers such as Proton Exchange Membrane Water Electrolyzers (PEMWEs), Alkaline Water Electrolyzers (AWEs), and Anion Exchange Membrane Water Electrolyzers (AEMWEs) are known to be sensitive to water quality, with a range of common impurities impacting performance, hydrogen quality and device lifetime. Purification of feed water adds to cost, operational complexity, and design limitations, while failure of purification equipment can lead to degradation of electrolyser materials and components. Increased robustness to impurities will offer a route to longer device lifetimes and reduced operating costs but understanding of the impact of impurities and associated degradation mechanisms is currently limited. This critical review offers, for the first time, a comprehensive overview of relevant impurities in operating electrolyzers and their impact. Impurity sources, degradation mechanisms, characterisation techniques, water purification technologies and mitigation strategies are identified and discussed. The review generalises already reported mechanisms, proposes new mechanisms, and provides a framework for consideration of operational implications.

Received 31st October 2022
Accepted 13th February 2023

DOI: 10.1039/d2se01517j

rsc.li/sustainable-energy

1 Introduction

1.1 Background

Low temperature water electrolyzers are increasingly being deployed to produce green hydrogen worldwide.¹ With applications in transport, heating and the chemical industry, the

availability of cheap low carbon hydrogen will be crucial to maximise the use of renewable energy and reduce or eliminate net CO₂ emissions.^{2–4} Electrolyzers are currently being deployed at the MW scale⁵ but to make a substantial impact on climate change, cumulative deployments need to be on the TW scale. To facilitate this Hydrogen Europe's strategic innovation and research agenda has set targets for a reduction of the cost of hydrogen fuel from € 5–8 to <€ 3 kg_{H₂}⁻¹ by 2030. This necessitates technology improvements. Thankfully there are still many opportunities to improve the understanding of the

^aNational Physical Laboratory, Teddington, Middlesex, TW11 0LW, UK. E-mail: graham.smith@npl.co.uk

^bDepartment of Materials, Imperial College London, Royal School of Mines, London SW7 2AZ, UK. E-mail: i.stephens@imperial.ac.uk



Hans Becker received his BSc and MSc in Chemical Engineering from Institut Teknologi Bandung and the Hong Kong University of Science and Technology, respectively. In 2018, he obtained his PhD in Electrochemistry from the Technical University of Denmark working on high temperature – Proton Exchange Membrane (PEM) fuel cells. Hans then moved to the National Physical Laboratory

UK for 3 years as a Higher Research Scientist working on the metrology of PEM fuel cells and electrolyzers. Hans is currently a Green Hydrogen Scientist at bp, leading the research of various water electrolyser technologies.



James Murawski received his MSci in Chemistry with industrial placement from the University of Southampton in 2018. He is currently pursuing a PhD under the supervision of Dr Ifan E. L. Stephens alongside industrial supervisors from Johnson Matthey and National Physical Laboratory. James' research focuses on benchmarking and improving understanding of oxygen evolution

catalyst stability in both aqueous model systems and full cell testing for PEM water electrolyser applications.



processes operating in electrolyzers and use this understanding to improve electrolyser lifetime and performance.^{6–8} One little-studied area of practical importance is in understanding the impact of water impurities on the performance and durability of electrolyzers.

Electrolysers producing hydrogen require water to operate; approximately 45 kg of water is electrolysed to produce 5 kg (~166 kW h_{HHV}) of hydrogen, enough to fill the storage tank in a typical passenger fuel cell electric vehicle. In comparison, around 5 kg of fresh water is used in thermoelectric power plants to produce 1 kW h of energy,⁹ while the typical UK citizen has a domestic water consumption of 142 kg per day¹⁰ and a total daily water footprint estimated to be in excess of 4600 kg.¹¹ Rather than quantity, the key challenge is the quality of the water.

It's well known that the water supplied to the electrolyser must be of high purity. Commercial electrolyser manufacturers

typically specify a minimum required water supply quality in terms of conductivity, typically <1 μS cm⁻¹ (>1 MΩ cm),¹² and a total organic carbon (TOC) content. Academic research groups typically use ultra-pure, ASTM D1193-06 Type I water (resistivity 18.2 MΩ cm @ 25 °C, TOC < 50 μg L⁻¹) supplied from a lab system, but the use of ion exchange resins or procedures to maintain resistivity in recirculating systems varies.¹³ The latest EU low temperature water electrolyser testing guidelines from the JRC¹⁴ call for ISO 3696 Grade 2 water, which has a conductivity of <1.0 μS cm⁻¹ at the inlet and for PEM electrolysis also require the anode water recirculation loop to have ion exchange resins and a conductivity meter in the loop.

Deionisation can be a large burden on the cost of producing green hydrogen. In a recent analysis, deionised water purification comprised 22% of the total balance of plant (BoP) cost of a 1 MW PEMWE system and was relatively insensitive to both system size and annual production rate.¹⁵ Deionisation also



Dipak completed his PhD at Hanyang University (Seoul, South Korea). His doctoral research involved the synthesis and characterisation of metal oxide and chalcogenide nanostructures for use in solar cells and water splitting devices. His subsequent postdoctoral work included roles at POSTECH (Pohang, South Korea), working on perovskite solar cell fabrication, and at the Italian Institute

of Technology (IIT, Genoa), where he worked to develop electrochemical experiments to investigate the hydrogen evolution and CO₂ reduction reactions. Dipak's current work at NPL involves the development of new characterisation techniques for water electrolyzers to understand their performance degradation.



Gareth is an NPL Fellow and Science Area Leader in the Electrochemistry Group at the National Physical Laboratory in Teddington, United Kingdom. His primary expertise is in the development of novel in situ diagnostic techniques and standard test methods for assessment of corrosion and material degradation in energy applications. Gareth is a fellow of the Royal Academy of Engineering

and holds visiting academic positions at UCL, the University of Strathclyde and Harbin Institute of Technology. He is the author of over 200 publications and is currently President of the World Corrosion Organization and Vice President of the European Federation of Corrosion.



Ifan is a Reader in Electrochemistry at the Department of Materials, Imperial College London. After a PhD at the University of Cambridge, Ifan moved to the Department of Physics at the Technical University of Denmark as a post-doctoral researcher, then Assistant Professor and finally as an Associate Professor. In 2015, Massachusetts Institute of Technology awarded Ifan the

Peabody Visiting Associate Professorship. Ifan has published 80 papers on topics including oxygen reduction, oxygen evolution, CO₂ reduction and nitrogen reduction. Ifan co-founded HPNow, focused on H₂O₂ electrosynthesis. Ifan and colleagues were awarded the John Jeyes Award for their work on H₂O₂ production.



Graham leads the UK's National Physical Laboratory's work on electrochemical hydrogen technologies and more broadly coordinates NPL's 'Metrology for Clean Hydrogen Energy' activities. His research focuses on developing high-quality measurements to support the global industrial and academic effort towards better electrochemical energy technologies. After an MSci and PhD at Imperial

College London, Graham carried out post-doctoral research at the University of Southampton and the University of Cape Town. Prior to joining NPL, Graham worked as a researcher in the SINTEF (Norway).



incurs operational costs from energy use, water supply and maintenance. It is difficult to find reliable numbers for energy use associated with the production of high purity water but it is likely on the order of 5 kW h m^{-3} ,^{16,17} or 0.2 kW for 5 kg of H_2 . The energy demand for deionisation therefore likely represents less than 0.2% of the energy stored in the product hydrogen. As with any plant, the capacity influences the cost. For a 200 m^3 per day throughput reverse osmosis (RO) plant, a capital expenditure (CAPEX) of $\$0.5 \text{ M}$ and operating expenditure (OPEX) of $\sim \$3 \text{ m}^{-3}$ has been estimated.¹⁸ The cost shifts to $\$2 \text{ M}$ CAPEX and $\$2 \text{ m}^{-3}$ OPEX when the capacity is increased to 1200 m^3 per day. It should be noted that reports in the literature often assume the feed water of RO plants to either be seawater or brackish water. When the inlet of the RO plant is much cleaner, *i.e.*, freshwater, the CAPEX and OPEX may be lower than these assumptions.

More subtly, the need for high purity water limits electrolyser design flexibility. For instance, in PEMWE systems, high purity feed water is typically deionised further using ion exchange resins as it recirculates through the system, maintaining purity during operation. The temperature sensitivity of these ion exchange membranes effectively limits the recirculation loop temperature and intolerance to ions requires use of particular materials in the stack and BoP. The conservatism in design caused by these issues increases the cost of green hydrogen.

There are a number of different mechanisms by which water impurities may conceivably impact the normal operation, performance, or lifetime of electrolysers. Many of these have not

been directly observed experimentally in operating electrolysers, though there is evidence of their impact in analogous fuel cell systems or *in situ* experiments on relevant materials. Hydrogen Europe's Strategic Research and Innovation Agenda has set a 2030 target of 0.1–0.12% degradation per 1000 h of electrolyser operation.¹⁹ Therefore, even slow, and seemingly insignificant degradation processes can have a large impact on the viability of an electrolyser.

1.2 Structure and operation of low temperature electrolysers

There are three main categories of low temperature electrolyser, which are distinguished mainly by the type of electrolyte they use. The principles of operation of a PEMWE, AEMWE, and AWE are shown schematically in Fig. 1.

1.2.1 PEMWE. PEMWE stacks are fed with low conductivity water. To achieve efficient operation, a thin ($100\text{--}200 \mu\text{m}$) proton exchange membrane (PEM), typically perfluorosulphonic acid (PFSA), separates the anode and cathode electrodes. PFSA membranes consist of a hydrophobic backbone of polytetrafluoroethylene (PTFE), a perfluoroethylene side-chain and a hydrophilic sulphonic acid head group. This results in high ionic conductivity and the physical separation of produced hydrogen and oxygen. PFSA ionomers are also used to provide ionic conductivity within the electrodes, which are typically made of highly active, acid-stable nanoparticulate platinum group metal (PGM) electrocatalysts with loadings of $\sim \text{mg cm}^{-2}$. Platinum and supported platinum catalysts are

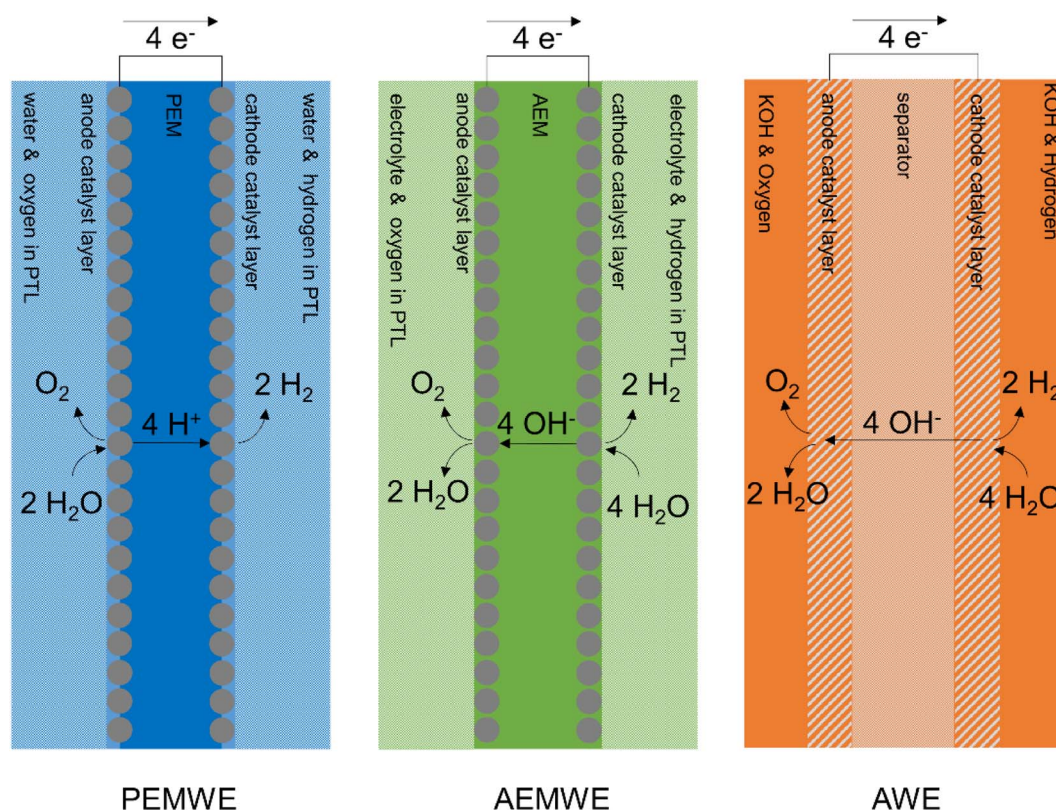


Fig. 1 Principles of operation of different low temperature water electrolyser technologies.



common at the cathode to catalyse the hydrogen evolution reaction (HER) with iridium oxide or mixed-metal oxides typically used at the anode to catalyse the oxygen evolution reaction (OER).^{20,21} Porous transport layers (PTLs) which support electrodes and manage two phase flow are usually made of titanium at the higher potential anode and sometimes carbon at the lower potential cathode. Bipolar plates (BPPs) which separate adjacent anodes and cathodes are usually titanium. PTL and BPP components are often coated with a thin layer of platinum in order to reduce contact resistance.⁶

PEMWEs are a semi-mature technology.^{6,7} They are typically operated with inlet water temperatures of around 60 °C and typical current densities of $\sim 2\text{--}4\text{ A cm}^{-2}$ resulting in acceptable cell voltages of 1.8 V and lifetimes of 6–9 years.²² Nonetheless, there is intensive research underway to increase lifetime, improve performance and decrease cost and these numbers are expected to improve. Operationally PEMWEs have a small volumetric footprint, are efficient, produce high purity hydrogen, can directly produce pressurised hydrogen and can rapidly respond to load variations which makes them well suited to handle the intermittency of renewable power.²³

1.2.2 AEMWE. AEMWEs have a similar structure to PEMWEs; however, as they are still primarily at an earlier stage of development there is a wider variation in the state-of-the-art.²⁴ Generally, they use an anion exchange membrane (AEM) to separate the anode and cathode and an anionic ionomer to provide conductivity in the catalyst layer. Typical AEMs are hydrocarbon polymers with pendant quaternary ammonium groups but there are a range of AEM chemistries and readers are referred to specialist reviews for further details.^{24–26} In some cases, AEMWEs are fed with a liquid electrolyte of ($<1\text{ M}$) aqueous potassium hydroxide or aqueous potassium carbonate.²⁵ The dominant ionic species is therefore hydroxide. The high pH allows the use of a range of nanoparticulate transition metal based catalysts, typically based on nickel or nickel-containing materials, at both the anode and cathode, though new catalysts are still under development and PGMs are still often used.²⁴

AEMWEs are of interest as they potentially retain many of the operational benefits of PEMWEs while avoiding the use of expensive and rare critical raw materials such as iridium and platinum. The latest AEMWEs in the academic literature tend to operate around 60 °C at 2 A cm^{-2} and $\sim 1.9\text{ V vs. RHE}$.^{27,28} Like PEMWEs, they generally have fast response times, produce pure hydrogen, have a small volumetric footprint, and directly produce pressurised hydrogen. Current drawbacks include the comparatively slow rate of hydrogen evolution reactions under alkaline conditions²⁹ and lifetime limitations (<4 years); with membrane, ionomer and catalyst developments needed in order to meet the long lifetimes required of commercial systems.³⁰

1.2.3 AWE. AWEs use aqueous solutions of potassium hydroxide, usually at concentrations of 20–50 wt%, to provide ionic conductivity between the anode and cathode electrodes. A porous, chemically inert separator such as Zirfon™, a composite of zirconia (ZrO_2) and polysulphone, is used to separate the evolved gases while supplying pathways for the

potassium hydroxide to provide ionic conductivity.³¹ As in AEMWEs, a range of transition metals are stable, and electrodes have often been made of porous bulk nickel with coatings such as RANEY®–Nickel to increase the electrode surface area.

AWEs are very mature systems and have been used industrially for almost a century, though the recent interest in electrolysis for energy applications is driving advances, for example zero-gap configuration cells.³² Typical current densities are 1 A cm^{-2} at 1.8 V, which results in larger footprints than membrane-based electrolyser. A 1 GW facility using PEMWE was estimated to have a footprint of 8–13 ha compared to AWE with 10–17 ha.³³ The liquid electrolyte and high permeability of the porous separator to gases hinders operation at very low currents, which makes coupling with renewables more challenging and increases the difficulty of producing directly compressed gas. However, AWE systems have a low capital cost compared to PEMWE and are very robust, with examples operating for 30–40 years.³⁴

1.2.4 Balance of plant. The stack is only one part of an operating electrolyser; in order to function, BoP is required. This generally refers to the sub-systems and components required to supply and recirculate water to the electrodes, manage heat, separate out and purify product hydrogen and oxygen and connect the electrolyser stack to the power source. In many cases the engineering of the BoP is limited by the stack operational parameters and will in turn strongly influence stack lifetime and performance.

1.3 Structure of the review

This review summarises existing published work on the behaviour of impurities inside low temperature electrolysers, highlights possible poisoning mechanisms based on *ex situ* and fuel cell experiments and identifies critical areas for future research. It also summarises techniques to measure the presence of impurities *in situ*, potential methods to improve tolerance to water impurities and strategies to recover performance after contamination events. Throughout, this paper draws upon a thorough review of the relevant electrolyser literature and is also informed by relevant results in fields such as proton exchange membrane fuel cells (PEMFCs) where the study of the impact of common impurities is more complete.

Section 2 reviews the potential sources of impurity for the different electrolyser technologies, covering both exogenous impurities in the water or electrolyte and endogenous impurities arising from the degradation of stack and ancillary components. Section 3 addresses the impact of specific contaminants on PEMWEs, and Section 4 does the same for AWEs and AEMWEs. Section 5 provides an overview of the available characterisation techniques to measure the impact of impurities during *in situ* or *ex situ* measurements, together with a summary of commonly-used water purification technologies and a discussion of possible mitigation and recovery strategies. The review concludes with a summary and an outlook for the study of the impact of impurities in electrolysers.



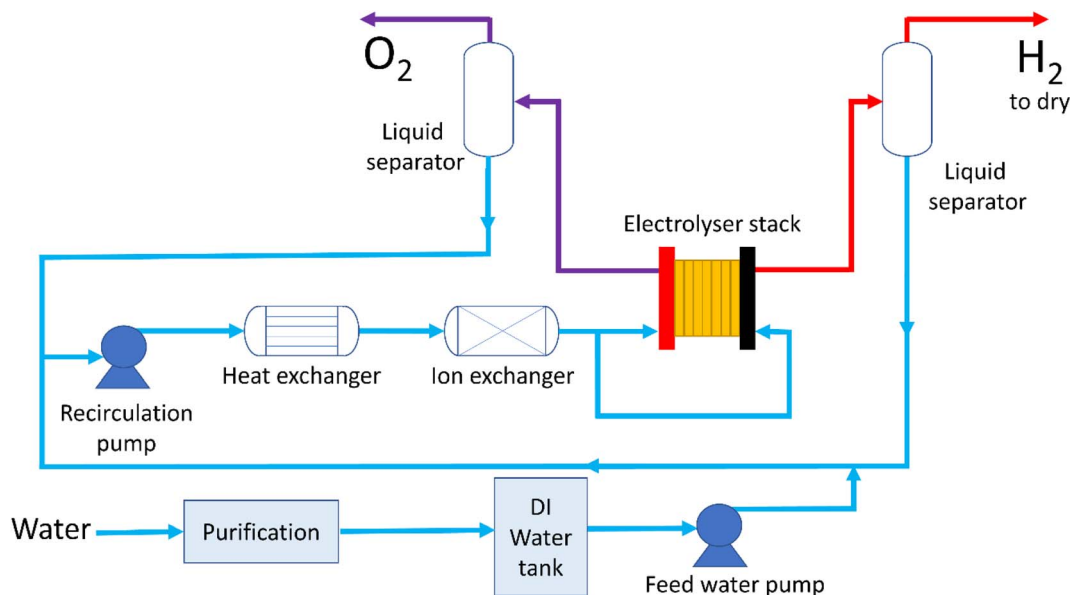


Fig. 2 Schematic of typical water process flow for a PEMWE stack with water flowing to both anode and cathode.

2 Sources of impurities

The water supply process in an electrolyser system is fairly complicated, with the BoP conditioning the water both before the inlet and in the recirculation loop, Fig. 2. Specific commercial configurations are proprietary but the scheme in Fig. 2 presents an illustrative example for the case where water is supplied to both anode and cathode. Water is sourced, usually from a local water supply, and purified in a multi-step process, details of which are often proprietary to the system manufacturer. Water is then fed to the stack where it is electrolysed to produce hydrogen and oxygen; it's usual to operate liquid flows far in excess of that required to sustain the electrochemical reaction to ensure the stack operates at high efficiency and to control the temperature. The temperature of the feed to the electrolyser depends on the specific technology but inlet temperatures of 60 °C to 80 °C are typical. As the electrolyser technologies discussed here operate exothermally, the vast majority of the thermal energy is lost by heating the water. Liquid and product gases usually exit the stack together and undergo some separation process with the liquid being recirculated and the gases exiting the system. To achieve very high purity hydrogen gas (with <5 ppm of water³⁵) there are further drying processes which are not considered here.

The recirculated water (or in the case of alkaline systems aqueous potassium hydroxide) must be cooled before supply back into the cell, though again the specific processes are usually proprietary. For PEMWE systems, ion exchange resins are used to remove contaminating anions and cations from the water supply and conductivity probes are commonly used to check for the build-up of ionic impurities. It is not possible to use ion exchange resins with alkaline electrolysers, due to the high concentration of potassium hydroxide. Behaviour of water in the stack is fairly intricate. In PEMWE systems it's common

to feed only to the anode where the water is consumed by the electrochemical reaction so that the only water in the cathode outlet is that which crosses through the membrane owing to diffusion and electro-osmotic drag. The same mechanisms operate in alkaline cells with the added complexity that water is consumed on the cathode and produced at the anode stoichiometrically. In alkaline cells using a supporting electrolyte there is also the need to rebalance the concentration of the electrolyte, which will vary over time.³⁶

Based on the process described above, it's possible to identify two different sources of impurities: exogenous and endogenous. We define exogenous impurities as those introduced by the feed water while endogenous impurities arise from degradation processes within the electrolyser and BoP itself. The following sections will discuss in detail the possible sources of impurities but with limited quantitative information available in the public domain from real electrolyser operation there remains much uncertainty.

2.1 Exogenous impurities

Water is an excellent solvent for cations and anions. Electrolysers typically require water that has been purified to a high resistivity, *i.e.*, featuring a low concentration of charge-carrying ionic species. However, in reality, impurities may still enter an electrolyser. Table 1 compares three types of water and the concentration of ionic impurities: seawater, tap water, and ASTM Type II water (1 MΩ cm). Tap water and seawater are widely used feedstocks for water electrolysis, which need to be purified to the required specifications of ASTM Type II water before use.

Compliant ASTM Type II water may still contain 5 μg L⁻¹ Na⁺ and 5 μg L⁻¹ Cl⁻, with the presence of other ions resulting in conductivity up to 1.0 μS cm⁻¹. Crudely then, a 1 MW PEMWE stack operating at full capacity for 10 years may therefore be



Table 1 Typical ionic impurity concentrations in different types of water^{13,37,38}

Ion	Seawater conc./mg L ⁻¹ (ref. 37)	Tap water conc./mg L ⁻¹ (ref. 38)	ASTM Type II water/ $\mu\text{g L}^{-1}$ (ref. 13)
Na ⁺	11 000	62	≤0.005
Ca ²⁺	400	51	—
Mg ²⁺	130	7	—
Al ³⁺	—	0.004	—
K ⁺	400	—	—
Cl ⁻	200	79	≤0.005
Br ⁻	10	—	—
F ⁻	1	—	—
HCO ₃ ⁻	110	158	—
CO ₃ ⁻	20	—	—
SO ₄ ²⁻	2800	47	—
NO ₃ ⁻	—	0.82	—

exposed to over 40 g of Na⁺, enough to ion exchange approximately one third of the membrane in such an electrolyser.† It is also important to note that, if not removed, impurities will become enriched; while water is consumed stoichiometrically in the electrolyser, the impurities it carries may not be. This process leads to a gradual increase in concentration of impurities in the cells and water recirculation loop, potentially by many orders of magnitude. Together, these issues justify the use of ion exchange resins in the recirculation loop of PEMWEs and clearly illustrate the possible implications of even a seemingly insignificant concentration of impurities. As mentioned previously, in-line ion exchange resins are not suitable for AWEs or AEMWEs operating with supporting electrolyte as the resins would quickly become saturated with the cations and anions from the electrolyte.

Beyond ions, other impurities may enter the electrolyser *via* the water supply. Organic impurities do not necessarily contribute to conductivity and the ASTM Type II water standard permits the presence of 50 $\mu\text{g L}^{-1}$ TOC. Biological components are also present in many water sources, and care must be taken to periodically clean pure water reservoirs to prevent their growth. Dissolved gases such as nitrogen, oxygen, argon, carbon dioxide, *etc.* will also be present in concentrations from nmol L⁻¹ to $\mu\text{mol L}^{-1}$.³⁹ While chemically inert gases are not likely to impact electrolyser performance, they may affect hydrogen quality, which is often required to meet the stringent quality standards.⁴⁰ Oxygen and carbon dioxide, on the other hand, are not wholly inert, as discussed below.

While most impurities may be removed during purification, accidental water contamination due to equipment or maintenance failures must also be considered. A 2016 presentation by Proton Onsite (now NEL Hydrogen) showed that in the period 2006–2015, 65% of stack defects were caused by inadvertent customer contamination⁴¹ while errors in system design and maintenance may also lead to the introduction of impurities. Due to the cost associated with purification, and in many cases local scarcity of groundwater or potable water, there is growing academic interest in developing electrolysers capable of

operating on lower quality water, or even on humidity from ambient air.⁴² Development towards cells able to run on seawater was the subject of a recent review.³⁷ Generating hydrogen by the electrolysis of small organic compounds or their aqueous solutions is also of academic interest and has been the subject of recent reviews.^{43,44}

2.2 Endogenous impurities

We classify any impurities arising from inside of the electrolyser system as *endogenous*. Operation of electrolysers for decades will lead to degradation of both the stack and BoP components, generating impurities *in situ*. Leaching from components⁴⁵ and contamination from manufacturing, commissioning and maintenance may also result in impurities being introduced by the electrolyser itself.

Much of the BoP, including tubes, tanks, gaskets, pumps, heat exchangers, sensors, filters, *etc.* are necessarily in intimate contact with the water supply to and from the electrolyser stack, which is both heated and saturated with either oxygen or hydrogen. Any corrosion products or leachants from BoP components are therefore able to enter the water supply. As there is little information available on the composition of water in operating commercial electrolysers it is difficult to quantify the likelihood and concentration of these impurities but there is literature evidence to suggest their presence at ppm concentrations.⁴⁵ Twenty different components commonly used as sealants in PEMFCs were leach tested in 90 °C DI water and showed tens to hundreds ppm of leachant.⁴⁵ There are reports of metals, even high-grade stainless steels, undergoing corrosion processes in ultrapure water with the expectation that ions are released,⁴⁶ though we stress there is a paucity of good quality data under conditions directly applicable to electrolysers. Such corrosion processes may be accelerated when ions are present in the electrolyte, either due to the build-up of impurities from other sources or because of the potassium hydroxide in AWEs or supporting electrolyte in AEMWEs. An example of contamination in PEMFCs attributed to ions leaching from tubing was reported by Sun *et al.* The performance of a 9 cell stack operated for 7800 h was recovered almost completely by immersing the catalyst coated membrane (CCM) in 0.5 M sulphuric and subsequent washing in DI water⁴⁷ with *post mortem* analysis of

† Assuming an electrolyser using Nafion-115 operating constantly at a current of 2 A cm⁻² and a water stoichiometry of 1.



the CCM revealing copper, calcium, and iron within the membrane and the catalyst layer. Grigoriev *et al.* conducted a 5500 h PEMWE experiment consisting of on/off current cycles to simulate intermittent renewable energy operation, albeit with platinum as the anode.⁴⁸ Unsurprisingly, the platinum was found to corrode and deposit within the membrane, with membrane thinning reported, but traces of silicon, potassium, and calcium contamination within the membrane were also reported. There were no traces of iron, nickel, or copper; therefore, it was concluded that impurities arose from degradation of water quality, with silicon suspected to leach out from a polypropylene tank into the water.

With the potential of the anode electrode routinely exceeding potentials of 1.5 V *vs.* RHE,⁴⁹ and with parts of the cell either very highly acidic or basic, materials for components inside the stack must be carefully selected to avoid corrosion and strike a balance between cost, performance and lifetime. The typical route taken has been to choose materials with very high stability such as platinum-coated titanium for PTLs and BPPs in PEMWEs and nickel for components in AWEs. In AWEs, there is an absence of literature for nickel or iron-based bipolar plate degradation, and studies of these materials are often performed under the scope of catalysis research. PEMWE components, on the other hand, have been shown to introduce impurities even when expected to be stable.

When an uncoated titanium PTL was used at the anode for 1000 h, titanium corrosion was reported with titanium oxides found deposited at the cathode.^{50,51} This phenomenon took place during both constant current and current cycling operation. Similar titanium contamination was also reported when a titanium alloy with 1 wt% of iron was used as the anode PTL.⁵² After 4000 h of operation, up to 7 wt% titanium and 3.5 wt% of iron was detected in the cathode catalyst layer, sourced from migrated corroded titanium. The authors suggested that this must primarily stem from the cathode titanium PTL, where a sufficiently protective titanium dioxide layer does not form due to the very acidic and reductive environment. A half-cell experiment also showed that both titanium and iridium dissolved into the electrolyte solution when the operation mode was switched from OCV to 1 or 2 mA cm⁻².⁵³ Despite the relatively low current density and operation in a specially designed cell, this demonstrates that dissolution can occur during transition from open circuit to another current. Siracusano *et al.* also performed a 3500–5700 h durability test at 1 A cm⁻² and 55 °C for a single cell PEMWE using CCMs manufactured *via* different hot-pressing protocols.⁵⁴ They reported titanium and iron impurities present in the anode catalyst layer, while no impurities were detected at the cathode. The authors proposed that iron likely comes from stainless steel tubing degradation, while titanium was suspected to be corroded due to hydrogen fluoride formation. The hydrogen peroxide generated from oxygen crossover at the cathode may also trigger titanium dissolution, although the exact mechanism is not clear.⁵⁵ There is a pressure to move from very stable materials such as titanium to less expensive materials. Though there is some scope to relax material requirements, for instance by recognising that not all parts of a PEMWE anode experience very high

potentials,⁵⁶ this results in an increased risk of components releasing ions and impacting the cell performance. Taken to the extreme, an uncoated stainless steel component was used as the PTL at the anode of a PEMWE^{57,58} and, unsurprisingly, extensive corrosion was observed within 15 h, and iron and nickel were detected at the anode, membrane and cathode.

Catalysts for electrochemical processes are chosen based on their stability, performance, and cost. With PEMWE catalysts for OER regularly showing an intrinsically inverse relationship between activity and stability,⁵⁹ there is necessarily a trade-off. For instance, ruthenium oxides are well known to be excellent catalysts for OER in acid but iridium oxides (or a mixed oxide based on iridium and ruthenium) are more commonly chosen for their stability.^{59,60} Even iridium oxides are not completely stable, undergoing electrochemical dissolution at operational potentials,⁶¹ with possible heightened dissolution rates during transient operation as the oxidation state may change.^{62–65} PEMWE cathode catalysts are also susceptible to corrosion, as the cathode can be exposed to oxidising potentials >0.9 V *vs.* RHE during shutdown periods.⁴⁹ Platinum black catalyst, for instance, was reported to lose 152 ng cm⁻² of a 3 mg cm⁻² loading over a 90 h OCV period.⁶⁶ While negligible, corresponding to only 0.005% of total cathode loading, long periods of operation can easily accumulate these ions in the cathode water. Catalyst supports are less used within electrolyser research than in fuel cells but their increased development and use offers both another vector for impurities to be introduced and also a potential mitigation, with supported fuel cell catalysts routinely showing reduced sensitivity to impurities.⁶⁷

While iridium (and/or ruthenium) dissolved during operation can redeposit at the anode, the dissolved species have a driving force to cross the membrane where they may be reduced by hydrogen. Deposition on the cathode catalyst may result in performance losses, and deposition inside the membrane as small particles can weaken the structural integrity of the membrane.⁶⁸ *Post mortem* investigation *via* scanning transmission electron microscopy X-ray energy dispersive spectroscopy (STEM-XEDS) of a low loading MEA (anode: 0.08 mg_{Ir} cm⁻²) tested for 4500 h @ 1.8 A cm⁻² showed significant crossover of iridium as well as its distribution across the entire CCM, Fig. 3. In addition to the iridium crossover, dissolved platinum, thought to occur due to dissolution of smaller nanoparticles during start up and shutdown procedures, was demonstrated to redeposit in the membrane resulting in similar effects to iridium redeposition but to a lesser extent than the anode catalyst.

The high pH present in AWEs and AEMWEs means that non-precious metals such as nickel are more stable to dissolution, as shown from their Pourbaix diagrams⁶⁹ and therefore cobalt, nickel, and iron based catalysts are frequently used at both the anode and cathode and exhibit high stability.^{70,71} In contemporary commercial AWE systems, nickel-based catalysts are the standard^{25,31,72} with systems exhibiting stability for up to 30 000 h.⁷³ This degree of stability is not yet observed in AEMWE and there is as yet no standardised catalyst for this technology.⁷⁴ One study of an AWE cell with a 316L stainless steel anode and a RANEY[®]-Ni cathode exhibited a negligible change in iron and



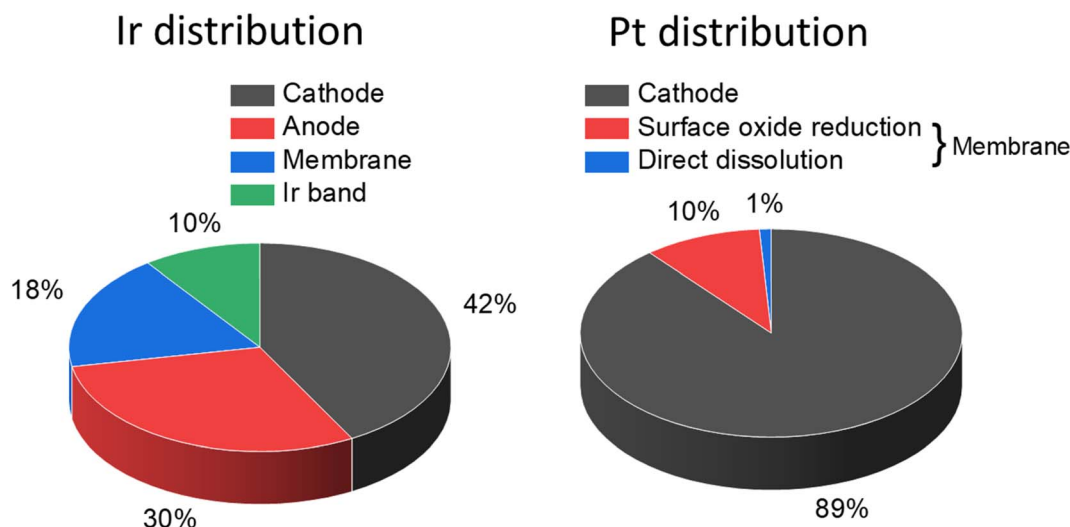


Fig. 3 Distribution of iridium and platinum across CCM after testing a low loading MEA (anode: $0.08 \text{ mg}_{\text{Ir}} \text{ cm}^{-2}$) for 4500 h @ 1.8 A cm^{-2} . This figure has been adapted from ref. 68.

nickel concentrations in the electrolyte after operation for 30 days at 0.3 A cm^{-2} in 6 M KOH at $75 \text{ }^\circ\text{C}$.⁷⁵ Another study using a specially devised ultraviolet-visible spectro-electrochemical cell reported that at potassium hydroxide concentrations of $0.1 \text{ M Ni(OH)}_2/\text{NiOOH}$ dissolves into the electrolyte at potentials of 1.2 V , 1.4 V , and 1.52 V vs. RHE . While the exact ionic species is uncertain, the absorption bands suggest that Ni^{2+} ions are the predominant species.⁷⁶ PGM based catalysts have also been explored for AWEs and AEMWEs,^{77–79} although most research is limited to activity studies in 3-electrode setups.^{80–82}

Despite improved PFSA membrane chemistry, membrane degradation as a result of hydroxy radical attack initiated by the Fenton reaction can occur and may release anions into the water.²⁸ Degradation of the PFSA polymer releases SO_4^{2-} from the end group and F^- from the polymer main group and side chain as a by-product.⁸³ Anion exchange membranes and ionomers are also subject to rapid degradation and may form various by-products.^{26,84} Indeed, the phenyl groups often found in hydrocarbon-based ionomers are known to adsorb strongly on the surface of AEMWE catalysts.³⁰ In PEMFCs, additives such as ceria or manganese oxides that are sometimes added to membranes and catalyst layers to act as radical scavengers have been shown to be redeposited throughout fuel cell electrodes^{85,86} and to migrate into the exhaust water;⁸⁷ similar behaviour may be expected with these materials in electrolyzers.

Impurities may also originate from residual chemicals associated with the manufacture of components within the electrolyser system. In AWEs, potassium hydroxide is added to the water initially (note it is not consumed in the reaction so the system only requires a pure water supply) and bulk potassium hydroxide is routinely contaminated with ppm levels of impurities such as Cl^- , PO_4^{3-} , SO_4^{2-} and a range of metals including aluminium, arsenic, calcium, copper, iron, sodium, lead, zinc, and silver ions.⁸⁸ Residual solvents or salts can also be present from the manufacturing of catalysts and membranes. For instance, chloride-based precursors of platinum and iridium

oxide nanoparticles can leave chlorine residues on the catalyst,^{89,90} while residual alcohol solvents such as methanol, ethanol, or propanol may result from the use of inks and slurries. Membranes also contain some residual impurities from the production process, which is why many research groups pre-clean purchased CCMs by immersing them in hot hydrogen peroxide and sulphuric acid before use. The break-in period for PEMFCs is frequently attributed to the *in situ* cleaning of catalyst layers and membranes^{91–93} and such break-in periods are also observed in PEMWEs.²⁷

Finally, by-products from the electrolysis process and recombination reactions between hydrogen and oxygen may form highly reactive radical and peroxide species stoichiometrically.^{84,94} Though they have a low concentration and short lifetime they may eventually result in damage to components.

3 Impact of impurities in PEMWEs

3.1 Introduction

This section addresses the impact of endogenous and exogenous impurities at both the anode and cathode of PEMWEs. Impurities are classified into three main types: cationic, anionic, and organic (inert compounds are only briefly considered) with plausible mechanisms of operation and impact assessed as summarised in Fig. 4. For many classes of impurity there is limited quantitative information available for specific impurities, so here the discussion remains largely qualitative.

3.2 Cationic impurities

For PEMWEs, cations are by far the most problematic class of impurity owing to their ability to impact the catalyst, ionomer and membrane, and the number of metallic components in the system that present endogenous sources. Cationic impurities can conceivably degrade performance and lifetime *via* several



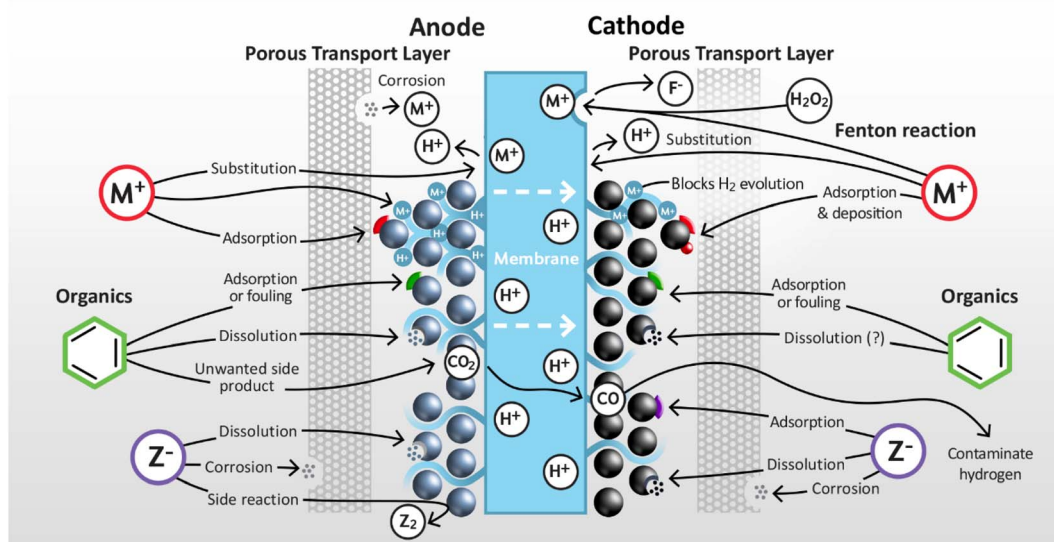


Fig. 4 Schematic illustration of the impact of various types of impurities on PEMWEs.

mechanisms: substitution of available protons in the membrane and ionomer, adsorption and deposition, and the initiation of unfavourable reactions.

3.2.1 Substitution

3.2.1.1 Membrane. Cations degrade PEMWE performance primarily by lowering the membrane conductivity.^{95–104} PFSA-based membranes such as Nafion possesses a very high conductivity in the presence of water due to the hyper acidity of the SO_3^- end group (estimated $\text{p}K_a$ of ca. -6) that results in the formation of mobile H^+ ions/protons.¹⁰⁵ However, most cations have a higher affinity than protons for the SO_3^- end group,¹⁰⁶ leading to proton displacement and a consequent reduction in the proton conductivity of the membrane. PFSA membranes in liquid water generally have a reported ionic conductivity in the range of $0.05\text{--}0.2\text{ S cm}^{-1}$,¹⁰⁵ yet this value can decrease by 1–2 orders of magnitude when all of the protons are exchanged to metal cations.¹⁰⁷

Not all impurity ions have the same impact; the situation is worse for higher valence ions which are preferentially absorbed, leading to higher concentrations in the membrane even at very low aqueous mole fractions (see Fig. 5).^{104,108–110} For example, Fe^{3+} has a $K_{\text{eq}} \sim 5000$ times higher than Na^+ . In some reports, at low concentrations (<10 ppm), absorption of Ni^{2+} and Cu^{2+} in the membrane was much higher than that of Fe^{3+} ; however, this was likely due to a relatively short membrane equilibrium time of 72 h.¹¹¹ There is a threshold concentration of ions before a step change in conductivity can be observed. Ions such as Na^+ , Cu^{2+} , Ni^{2+} , and Fe^{3+} were only observed to yield a significant decrease of conductivity at concentrations >10 ppm in water. The effect of these impurities could be mitigated slightly when present at low pH. Upon contamination in $1\text{ M H}_2\text{SO}_4$, cationic impurities such as Fe^{3+} or Cr^{2+} do not impose conductivity loss at concentrations below $200\text{--}300$ ppm.¹¹² On the other hand, contamination in distilled water results in conductivity loss at concentrations as low as 10 ppm.¹¹⁰

Non-metallic cations, such as ammonium (NH_4^+), can also substitute protons within the PFSA membrane.^{113–115} As with metallic cations, the membrane conductivity in PEMFCs has been reported to decrease with an increased concentration of NH_4^+ absorbed by the membrane in both liquid phase^{113,114} and gas phase.^{114,115} As PEMWE membranes are typically fully equilibrated with liquid water, the effect of ammonium contamination may be less severe than that seen in PEMFCs and where the loss in conductivity was found to be lower at higher relative humidity. Note also, that in PEMFC, catalyst poisoning may dominate performance losses introduced by ammonia.¹¹⁶

Apart from conductivity loss, cationic impurities can also compromise membrane water transport properties and overall membrane stability. Na^+ and Ca^{2+} adsorption has been reported

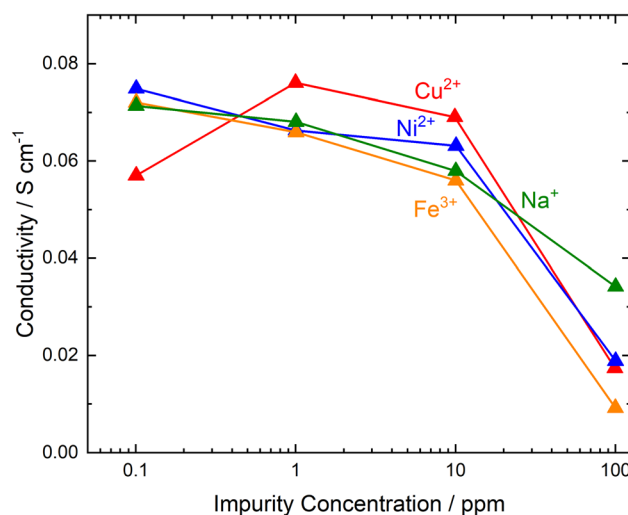


Fig. 5 Impact of immersion in cation solutions in distilled water for 48 h on PFSA conductivity. This figure has been adapted from ref. 110.



to change the water balance by increasing the water transfer coefficient ($t_{\text{H}_2\text{O}}$) and decreasing water content.^{95,96} The $t_{\text{H}_2\text{O}}$ for Nafion 115 increased from 2.5 to more than 10 as all the protons in the membrane were exchanged with Ca^{2+} .¹¹⁷ Compromised water drag is problematic for gaseous applications such as fuel cells though the situation is less clear for electrolyzers. Thermal stability is also affected by cations, with the onset temperature for thermal degradation of the SO_3^- group higher when Nafion® NR 211 membrane was contaminated with Ni^{2+} , beginning at a $\text{Ni}^{2+}/\text{H}^+$ ratio of 0.2.¹¹⁸

3.2.1.2 Ionomer. PEMWE catalyst layers routinely contain an ion conducting phase, referred to here as ionomer, to increase the electrochemically active surface area¹¹⁹ – the area where reactant (water), ion conducting phase (protons) and electrically conducting phase (catalyst surface) are in intimate contact. PEMWE ionomer is typically also a PFSA and therefore suffers from the same issues as the membrane when exposed to cations. While loss of protons in the ionomer may increase the observed resistance of the cell, the catalyst layers are thin (typically $<25\ \mu\text{m}$ (ref. 120)) with ionomer layers on the catalyst surface many orders of magnitude thinner than this. The effective loss of electrochemically active surface area that results as proton transport is deactivated is therefore generally a larger issue. This is confirmed by studies that have introduced metallic impurities directly into the water feed of operating PEMWEs. In one case, the impact on a PEMWE cell of incorporating 0.05 M Na_2SO_4 into the anode and cathode water feeds was studied with an *in situ* reference electrode.¹²¹ Following contamination, the pH of the anode water decreased from pH 6 to pH 3 as the protons in the anode ionomer and membrane were substituted and formed sulphuric acid in the water. Intriguingly, the pH of the cathode water increased from pH 6 to pH 11 after 3 h of operation. The increase of pH at the cathode was attributed to Na^+ migrating through the membrane and substituting for protons in the cathode catalyst layer, eventually resulting in a transition from the acidic HER to the alkaline HER, which releases OH^- . Interestingly, the effect depends on which water supply is contaminated. When the anode supply was contaminated the cathode potential decreased from 0 V to $-0.5\ \text{V}$ vs. RHE, while a much smaller decrease was observed when the cathode water was contaminated (0 V to $-0.1\ \text{V}$ vs. RHE). It's unclear whether the presence of the impurity impacted the operation of the *in situ* reference electrode, which can be highly sensitive to their local environment.

Electromigration of cations causes the cathode ionomer to be more severely impacted than the anode ionomer, particularly when operating at high current densities.^{122–126} Fig. 6 shows the effect when the anode of a PEMWE cell was poisoned with 0.001 M Na_2SO_4 at $0.5\ \text{A}\ \text{cm}^{-2}$; the anode overpotential plateaued then decreased as the cathode pH spiked to 11.¹²² A similar potential peaking trend was found in a study injecting 0.005 M $\text{Fe}_2(\text{SO}_4)_3$ with 0.01 M H_2SO_4 as the anode water feed of a PEMWE cell operated at $0.5\ \text{A}\ \text{cm}^{-2}$.¹²⁴ The phenomenon can be explained by cation electromigration to the cathode alleviating poisoning at the anode, while exacerbating it at the cathode. *Post mortem* Electron Probe Microanalysis (EPMA) showed cation accumulation in the cathode when the anode

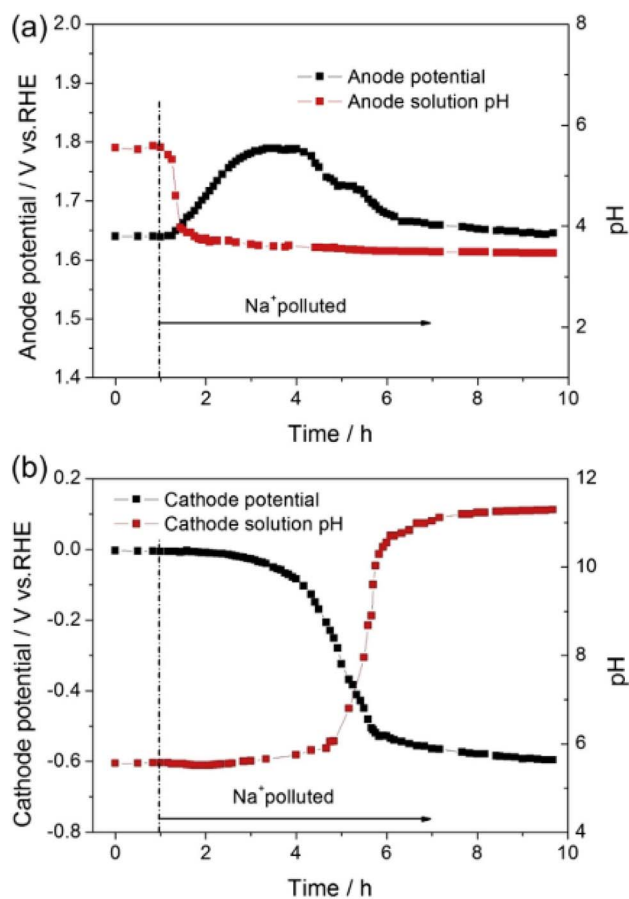


Fig. 6 (a) Anode potential vs. time and anode water pH vs. time; (b) cathode potential vs. time and cathode water pH vs. time for a PEMWE cell poisoned with 1 mM Na^+ at $60\ ^\circ\text{C}$, $500\ \text{mA}\ \text{cm}^{-2}$ with flow rate of $1\ \text{ml}\ \text{min}^{-1}$. This figure has been reproduced from ref. 122 with permission from Elsevier, Copyright 2013.

supply was poisoned by sodium¹⁰⁵ and iron.¹²⁴ Electrochemical Impedance Spectroscopy (EIS) also showed an increase in charge transfer resistance (R_{ct}) rather than ohmic resistance (R_{Ω}), indicating that the degradation mechanism is associated with loss of electrochemically active surface area. Damage to performance was also observed even at ppm levels of cationic impurities in water.^{125,126} A $\sim 0.2\ \text{V}$ increase in cell voltage at $2\ \text{A}\ \text{cm}^{-2}$ after 4 h was reported even with iron concentrations as low as 1 ppm, increasing to $\sim 0.9\ \text{V}$ at $2\ \text{A}\ \text{cm}^{-2}$ at 10 ppm,¹²⁵ with EIS indicating that an increase in kinetic overpotential seemed to be the major reason for performance degradation.¹²⁶

While most studies point to accumulation of cations in the cathode catalyst layer as the primary degradation mechanism, the governing mechanism is largely controlled by the rate of ion migration (a function of current density) balanced by ion back-diffusion.^{123,127} A study of trace ion impurities in the cathode effluent of a PEM electrolyser for hydrogen peroxide production showed that the concentrations of titanium and iridium cations generated by corrosion processes *in situ* were dependent upon the operating current density.¹²⁸ Gd^{3+} is a model trivalent cation used in neutron imaging studies due to its very high neutron cross-section, enabling good contrast resolution.^{123,127} Upon



passage of electrical current at 1 A cm^{-2} in a single cell PEMWE poisoned with Gd^{3+} , the ions accumulated in the cathode catalyst layer within $\sim 16 \text{ s}$. Once the current was switched off, accumulated Gd^{3+} back-diffused to the membrane within 200 s . This behaviour is identical to that reported for Fe^{3+} ,¹²³ and is therefore likely to be expected for most cation poisoning. In case of Fe^{3+} , voltage breakdowns at different current densities show that an increased R_{Ω} dominates at $<0.2 \text{ A cm}^{-2}$ but that increased mass transport and kinetic overpotentials dominate at $>1 \text{ A cm}^{-2}$.^{123,127} At low current densities, the rate of ion migration can be countered by Fickian back-diffusion arising from the concentration gradient between cathode and anode. This prevents excessive accumulation in the cathode and thus R_{Ω} degradation dominates at low current density. As the current density increases, back-diffusion is overpowered by electromigration, causing mass transport and kinetic overpotential to dominate degradation at high current density. Metallic cations are also much less mobile in the membrane meaning that they are not easily purged once present.

3.2.2 Deposition. Another way in which metallic cations can degrade PEMWE performance is *via* electrodeposition on the cathode,¹²⁹ where a potential in the range -100 mV to 0 V *vs.* RHE is routinely encountered. Metal cations with a more positive standard reduction potential than 0 V *vs.* RHE can be reduced to form a metal deposit on the surface of the cathode catalyst, leading to a decrease in accessible electrochemically active surface area. The potential window in which this can occur can be further enhanced by the process of underpotential deposition (UPD), particularly in the presence of chloride ions.¹³⁰

There are some reports where cations have been observed to deposit on the cathode during operation. A half-cell study of HER on platinum in the presence of 10^{-6} M CuSO_4 exhibited visible copper deposition.¹²⁹ The presence of CuSO_4 led to an increase to the steady state overpotential of 300 mV , which was reached more rapidly at higher concentrations ($<3 \text{ h}$ at $5 \times 10^{-4} \text{ M}$ *vs.* $>20 \text{ h}$ at 10^{-6} M). Another study investigated the impact of 0.01 M NiSO_4 in the feed water of a PEMWE cell operating with platinum on both cathode and anode.¹³¹ A decline in performance was observed when the cell voltage was held at 2.35 V (0.3 A cm^{-2} to *ca.* 0.15 A cm^{-2}) attributed to membrane conductivity decrease. Bulk nickel deposition was also suspected to take place; however, this is unlikely to occur in state-of-the-art PEMWEs as the cathode potential rarely drops below -100 mV *vs.* RHE.

PGM ions have also been shown to deposit within the membrane, creating local hotspots that promote membrane degradation. Cycling the catalyst potential up to 1.0 V and 1.2 V *vs.* RHE during operation in PEMFCs results in platinum dissolution into the membrane, where it is subsequently chemically reduced by crossover hydrogen, forming characteristic metallic bands.¹³² Membrane thickness also affects deposition rate, as thinner membranes promote hydrogen crossover and exhibit more platinum bands within the membrane post-potential cycling.¹³² A PEMWE with 0.08 mg cm^{-2} iridium oxide anode and 0.3 mg cm^{-2} Pt/C operated for 4000 h at 1.8 A cm^{-2} showed iridium and platinum migration to the opposite

electrodes,⁶⁸ as shown in Fig. 7. The majority of iridium (46% of total iridium) migrated and was deposited in the cathode, while 28% remained inside the membrane. Platinum migration was less severe, with only 18% of total catalyst deposited inside the membrane. Significant loss of membrane mechanical integrity was also reported after the test, but the correlation with the deposited platinum/iridium remains unclear.

Platinum deposition inside the membrane can have short-term beneficial side effects, as it can mitigate gas crossover, increasing hydrogen quality and ensuring that the flammability limit is not exceeded.^{133,134} Employing a thin platinum layer at 0.02 mg cm^{-2} loading between two Nafion 115 and 212 membranes was reported to reduce the concentration of hydrogen in oxygen from $0.2 \text{ vol}\%$ to $<0.005 \text{ vol}\%$ at 1 A cm^{-2} .¹³³ Even when the cathode was operated at 30 bar_a , 0.06 mg cm^{-2} platinum loading in a thin Nafion 212 membrane could suppress hydrogen in oxygen from $3 \text{ vol}\%$ to $<1 \text{ vol}\%$.¹³⁴ In all these studies, the reported downside is the notable increase of R_{Ω} due to the imperfect interface between the recombination catalyst and the membrane.^{133,134} However, the long-term adverse effects remain unknown as all reported tests in the literature were performed over short timescales.

3.2.3 Reactions. One specific impact of Fe^{2+} contamination is initiation of the Fenton reaction, the decomposition of peroxide to hydroxy radicals; radicals which may then attack the polymeric backbone of PFSA and are notorious for causing membrane thinning in PEMFC^{135,136} and PEMWE^{20,137} systems. Unlike in PEMFCs where membrane degradation occurs on both anode and cathode sides, multiple studies of membrane thinning in PEMWEs have confirmed that thinning takes place predominantly on the cathode side.^{94,138,139} At typical PEMWE cathode potentials ($<0 \text{ V}$ *vs.* RHE), crossover oxygen is predominantly reduced *via* a 2-electron process into hydrogen peroxide.¹⁴⁰ Membrane thinning was reported to be more severe at low current densities, where the rate of oxygen crossover is higher, and at elevated temperatures ($90 \text{ }^{\circ}\text{C}$) while accelerated cell degradation has been observed at low concentrations of Fe^{3+} ($1\text{--}10 \text{ ppm}$).^{125,126} Although results to date have focused on Fe^{2+} as a catalyst, other transition metal cations may also act as a catalyst for peroxide decomposition to hydroxy radicals.¹⁴¹

3.3 Anionic impurities

Unlike cations, anions cannot replace protons within the membrane and ionomer, and other mechanisms therefore operate. The most commonly highlighted problems are the initiation of side reactions, such as the chlorine evolution reaction that may lower hydrogen quality, and processes that accelerate the corrosion of metallic components. Literature reports predominantly feature investigations of chloride anions, and these therefore dominate the examples in this section, although it's likely that other anions have similar mechanisms of operation.

3.3.1 Reactions. The most common anionic impurities studied are halogen ions, with Cl^- being the most prevalent as it is present in by far the highest concentration in both seawater and potable water, Table 1. In the case of PEMWEs, Cl^- may



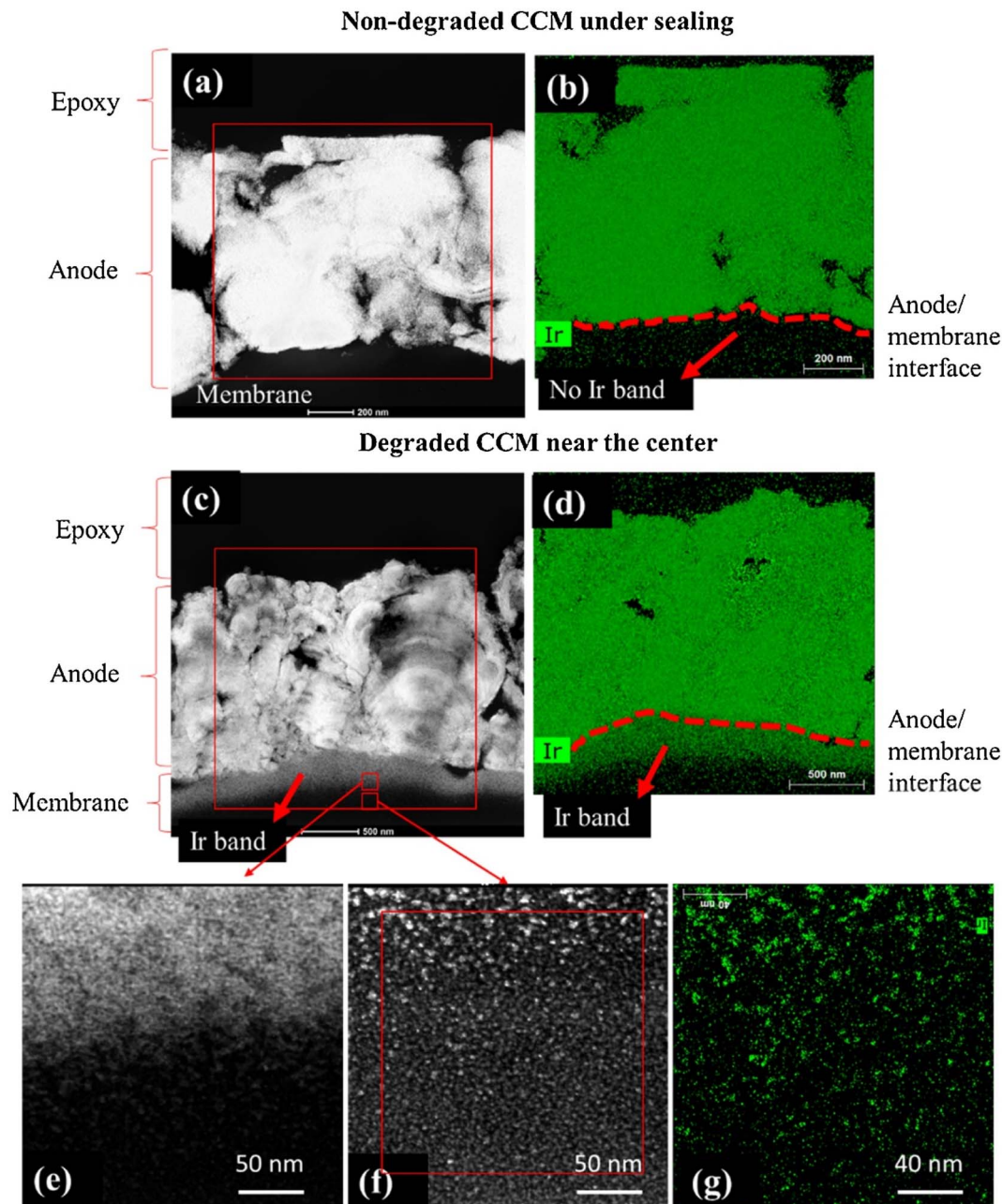
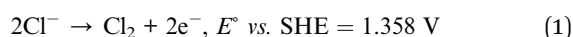
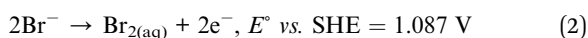


Fig. 7 Migration of Ir into the membrane (a, c) STEM images, (b, d) elemental mapping using STEM-EDS of non-degraded and degraded membranes (e and f) are high magnification images of Ir band while panel (g) shows elemental mapping of Ir for (f). This figure has been reproduced from ref. 68 with permission from Elsevier, Copyright 2020.

react *via* the chlorine evolution reaction (CER) that can compete with the OER at the anode, producing impurities in the hydrogen:



Other halide anions, such as F^- , Br^- , and I^- , can also be oxidised to their gaseous form with analogous evolution reactions:



OER is more thermodynamically favourable than CER under acidic conditions relevant to PEMWEs¹⁴² as shown in Fig. 8, however during typical operation the PEMWE anode potential is higher than the CER onset potential. The CER is also more kinetically facile than the OER as the latter requires 4 electrons and is thought to have 3 intermediates, while CER only requires 2 electrons and has a single intermediate.^{143,144} As such the ratio



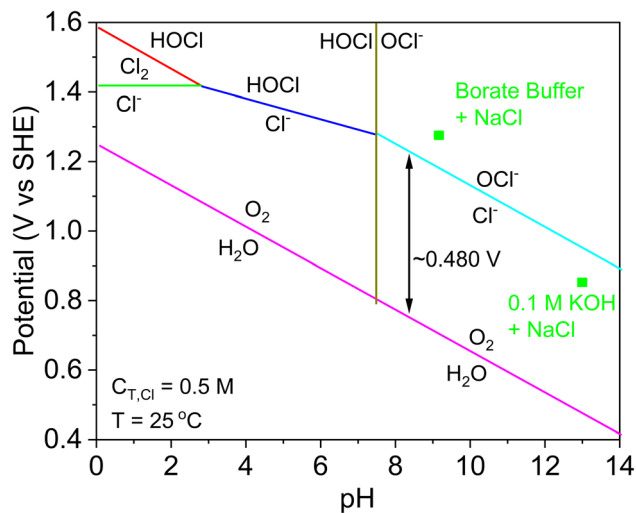


Fig. 8 Pourbaix diagram of the water-chloride electrolyte system. This figure has been adapted from ref. 148.

of exchange current density for CER/OER on most PEMWE anode materials ranges from 10^3 to 10^7 , and the evolution of chlorine is therefore much more favoured kinetically. For mixed oxide catalysts such as ruthenium dioxide, the selectivity for CER at $E = 1.25$ V (vs. Ag/AgCl) is >0.93 for compositions of ruthenium dioxide >20 wt%.¹⁴⁵ In fact, virtually any catalyst that is active for OER is also highly active for CER.¹⁴⁶ There is a growing trend of developing oxygen selective catalysts for use in saline conditions, but most of the work is conducted at alkaline pH.³⁷

A recent study of an iridium oxide catalyst on a rotating ring-disk electrode established that the OER and the CER reactions take place at different sites without competition;¹⁴⁷ prior to this study, the conventional wisdom was that the two reactions occupy the same sites. The oxygen production rate remained constant even when the Cl^- concentration was varied between 0–100 mM. The chlorine production rate increased linearly with

Cl^- concentration, with the CER selectivity approaching 100% at Cl^- concentrations >50 mM, Fig. 9.

3.3.2 Adsorption or fouling. Another mechanism whereby anions can pose a threat to cell performance is adsorption to catalyst surfaces and consequent reduction of the active surface area. This has been most prominently studied in the platinum catalyst system for PEMFC applications^{149–151} and redox flow batteries.¹⁵² Bromide ions, for instance, adsorb quite readily in the potential region of 0.5–1.0 V vs. RHE at 0.1 mM concentration,^{153,154} reaching complete monolayer coverage at 0.5 V vs. Ag/AgCl, 1 M KCl.¹⁵⁵ However, desorption of Br^- was observed in the hydrogen adsorption region of platinum, with complete desorption occurring at potentials lower than ca. 0 V vs. RHE.^{155,156} The concentration of Br^- also plays a significant role, as adsorption was found to be reversible when using 1 mM HBr, yet 3 M HBr induced an electrochemical surface area (ECSA) loss of $>50\%$ that persisted even at -30 mV vs. RHE.¹⁵⁷ Iodide, another halogen group anion, was found to adsorb as much as 0.4 monolayer on platinum at 0.2 V vs. Saturated Calomel Electrode (SCE).^{158,159} However, the steady-state concentration of adsorbed iodide was lowered when the electrode potential was held at -0.1 V vs. SCE. This hints at similar behaviour to Br^- , whereby if a sufficiently negative potential is applied I^- would likely desorb completely from the catalyst surface. Cl^- behaviour on platinum is similar to that of other halide anions. Adsorption at 1 mM has been reported to modify the charge associated with hydrogen underpotential deposition (H_{UPD})^{150,151} and at ca. 0 V vs. RHE, 0.1 monolayer of Cl_{ads} is still observed.¹⁶⁰

Overall, the strength of halide ion adsorption follows the order $\text{I}^- > \text{Br}^- > \text{Cl}^-$.¹⁶⁰ The platinum cathode of a PEMWE typically encounters a negative potential, in the range of -0.1 – 0 V vs. RHE.⁴⁹ Hence halide ions pose a risk by adsorbing on the catalyst surface and in the case of I^- and Br^- this may be irreversible at sufficiently high concentrations. HER on platinum in acid is, however, still a very fast reaction and so it stands to reason that minor anion contamination will not result in significant loss in performance. As a comparison, the hydrogen oxidation reaction (HOR) in PEMFCs only shows a major loss in performance when the catalyst is covered by 0.4–0.6 monolayer of CO.^{161,162}

Adsorption of chloride on the platinum on carbon cathode catalyst of PEMFCs has also been reported to enhance hydrogen peroxide production,^{150,163} which as mentioned above is a necessary precursor in the Fenton reaction that attacks the PFSA backbone. Adsorption of Cl^- accelerates the cleavage of the O–O bond, an important step in the hydrogen peroxide reduction reaction. Consequently, more hydrogen peroxide is released into the catalyst layer/membrane interface, promoting membrane thinning. In PEMWE cathodes, the low potential will likely hinder any hydrogen peroxide formation from taking place, though further study is needed to confirm this assumption.

Sulphate (SO_4^{2-}) and bisulphate (HSO_4^-) anions are other by-products of membrane degradation. SO_4^{2-} anions are well known to specifically adsorb onto platinum catalyst surfaces. SO_4^{2-} adsorption is a function of potential and on Pt(111) peaks

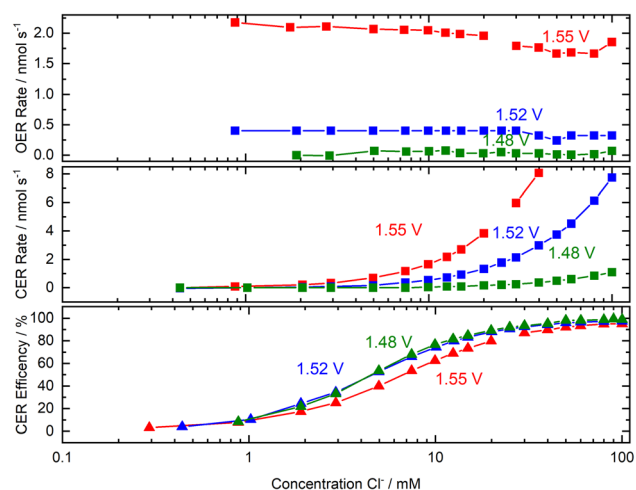


Fig. 9 Plot of OER and CER reaction rates as a function of $[\text{Cl}^-]$. This figure has been adapted from ref. 147.



in the range 0.2–0.8 V vs. Ag/AgCl, with almost complete desorption at –0.2 V vs. Ag/AgCl, have been reported.¹⁶⁴ At PEMWE cathode potentials (<0 V vs. RHE) and at the local acidic pH (from the Nafion ionomer), H⁺ adsorption overwhelms any SO₄²⁻ adsorption.¹⁶⁵ At the anode, SO₄²⁻ at high concentrations (0.05 M H₂SO₄) have been shown to adsorb on Ir(111) and shift the onset of the OER to higher potentials.¹⁶⁶ The lowest concentration of SO₄²⁻ adsorption on iridium catalysts studied to date, to the best of our knowledge, is 0.05 M H₂SO₄.¹⁶⁷ The concentration of SO₄²⁻ from internal membrane degradation would likely fall in the ppb range, which is unlikely to pose a significant threat.

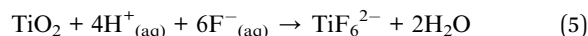
3.3.3 Corrosion of metallic components

3.3.3.1 Cathode catalyst. Another degradation mechanism associated with chloride anions is their tendency to form chloroplatinic ligands that enhance platinum dissolution,¹⁶⁸ a well-studied problem for platinum catalysts.^{169–177} The addition of just 2.8 μM of HCl to 1 M H₂SO₄ was reported to increase the dissolution rate of platinum by 2 orders of magnitude, even without any polarisation.¹⁶⁸ Under constant applied potential, platinum dissolution in the absence of Cl⁻ is slow; for example, potentiostatic testing of electrodes at 1.2 V vs. RHE in 0.5 M H₂SO₄ for 24 h did not exhibit any mass change as measured by electrochemical quartz crystal microbalance.¹⁷⁸ However, platinum is much more prone to dissolution when the potential is cycled between low and high potential, particularly during cathodic scans.^{179,180} Potential cycling from 0.0–0.8 V vs. SHE was reported to induce dissolution of platinum even without any Cl⁻ in the system, increasing substantially when the upper potential limit was 1.2 V; though we note the use of an Ag/AgCl reference electrode in the experiment, which could potentially leak trace amounts of Cl⁻ into the electrolyte.¹⁷³ Potential cycling of platinum takes place routinely in both PEMFCs and PEMWEs during start-up and shutdown.^{49,181} During shutdown of PEMWEs, the potential of the cathode may transition from <0.0 V vs. RHE to ca. 1.0 V vs. RHE due to crossover of oxygen from the anode. In the presence of Cl⁻ the rate of platinum dissolution during potential cycling is higher, and this has been investigated as a function of Cl⁻ concentration.^{174–176} The effect of ppm level Cl⁻ concentration on carbon supported platinum was studied using *ex situ* quartz crystal microbalance. The combined effect of dissolution and adsorption reduced the platinum ECSA by 4–13% when Cl⁻ concentration was varied between 500–2000 ppm.¹⁷⁶ The corrosion potential of platinum was found to decrease from 1.23 V to as low as 0.82 V when 0.01 M of Cl⁻ was present in the electrolyte.¹⁷⁴

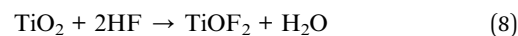
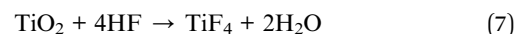
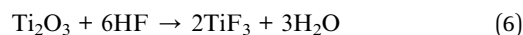
3.3.3.2 Anode catalyst. Unlike for platinum, there is a scarcity of studies on the impact of Cl⁻ on the stability of iridium oxide anode catalysts. Iridium in perovskite form (Ba₂BiRO₆) cycled triangularly from 1.6–0 V vs. RHE showed enhanced iridium dissolution in the presence of 50 mM NaCl. The chloride was reported to exclusively affect noble metal dissolution in the perovskites, which occurred during potentiodynamic control and active OER/CIER.¹⁸² Ruthenium was also shown to have enhanced dissolution in the presence of Cl⁻.¹⁷⁰ Platinum and ruthenium were cycled from 1.5–0.1 V vs. RHE in the presence of varying concentrations of Cl⁻,¹⁷⁰ and it was found

that in 0.1 M H⁺ + 0.01 M Cl⁻, the Ru dissolution was 11.8% after 120 cycles with dissolution increasing to almost 100% in 1 M Cl⁻. It should be noted that the oxide form of iridium and ruthenium is typically used at the anode, in which case the dissolution profile may be vastly different. For instance, a Dimensionally Stable Anode (DSA) that usually consists of iridium and ruthenium oxides is an industrial standard for the CER in the Chlor-Alkali Process and shows excellent stability in concentrated Cl⁻ solutions.¹⁸³

3.3.3.3 Porous transport layer/bipolar plate. Other components that can be attacked by anions are the PTL and BPP, which are typically manufactured from titanium, often with a platinum coating to minimise contact resistance. For platinum-coated components, the information presented above in the cathode section applies. For bare titanium components in oxygenated solution, an oxide layer forms at the surface of titanium and acts as a corrosion protection layer. This layer is even immune to the presence of Cl⁻ at 3.5% concentration.¹⁸⁴ However, anions such as F⁻ are reported to be able to attack the TiO₂ passive film, forming a soluble titanium-fluoride compound according to the following reaction:^{137,185}



A threshold concentration of F⁻ for titanium corrosion in the range 0.0005–0.02 M has been reported in 0.05 M H₂SO₄ and at 0.001 M F⁻ in 1 M HClO₄.^{185–187} Below the threshold concentration, the oxide film is partially damaged yet still protective. Above the threshold value, F⁻ converts the compact film to a porous non-protective layer.¹⁸⁵ Fluorine can also react with free protons to form HF, which is known to react with the surface oxide layer through the following reactions:



The rate of formation of HF depends on the proton concentration, *i.e.*, the pH of the solution. In solutions of 200 ppm and 400 ppm of F⁻, corresponding to a pH of 4.0 and 4.3 respectively, it was found that the critical HF concentration before the corrosion current shifted was around 21 ppm.¹⁸⁸ F⁻ ions are a common impurity in PEMWE water as they are generated *in situ* when the membrane degrades; measurement of F⁻ concentration is often used as an indicator of membrane degradation rate.¹³⁸ F⁻ has been largely overlooked as a potential corrodant in PEMWEs, and only a few studies have acknowledged its potential role in PTL and BPP degradation.^{54,137} However, at the typical pH of PEMWE water >1000 ppm of F⁻ may be required exceed the critical concentration.¹⁸⁸ When a PEMWE was operated at its normal conditions (60 °C, 2 A cm⁻²), <2 ppb of F⁻ was detected at the cathode outlet.²⁸ In order to trigger titanium corrosion, a commercial PEMWE with a correctly functioning water purification system would need to experience failure of both the ion exchange resin



and the conductivity probe. Even then, it would take a very long time for the F^- to accumulate to a concentration of 1000 ppm.

While chloride is typically the dominant ionic impurity in PEMWE feed water, the increase in concentration of any ions (anions or cations) can impact the water conductivity. In PEMWEs there is a distinct boundary between the highly ionically conducting catalyst layer and the very high resistance water circulating in the system. This means that the potential of electrolyser components such as the bipolar plate and large parts of a PTL are decoupled from that of the catalyst layer.⁵⁶ If the conductivity of the water rises, the extent of this decoupling decreases and may lead to more rapid corrosion of electrolyser components. This highlights the possible damage that even chemically inert ions may facilitate when accumulated to high concentrations.

3.4 Organic & inert impurities

3.4.1 Organic impurities. Organic impurities cover a wide range of species; in practice the specification of ASTM Type II water means that the presence of organic material in inlet water should not exceed 50 ppb on a carbon atom basis.¹³ However, as industrially relevant throughput systems, *e.g.* process waste streams, are implemented for electrolysis, the effect of organic impurities on electrolysers may become more important, both in (i) determining the quality of water purification required for standard PEMWE and AWE setups and (ii) as a result of direct seawater electrolysis where higher concentration of organics species may be present due to further constraints on purification stages.³⁷

There are two main avenues by which electrolyser performance could be impeded by organic contaminants in PEMWE systems: (i) poisoning *via* adsorption of species on the catalyst surface decreasing electrochemical activity and (ii) enhancing catalyst dissolution.¹⁸⁹ Non-ionic species are expected to have only a small impact on membrane and ionomer with PFSAs, well studied in analogous applications such as direct methanol fuel cells. In addition to directly poisoning the catalyst, the oxidation of organic molecules to carbon monoxide and carbon dioxide can also indirectly have a detrimental effect on the system. The ISO 14687 standard sets maximum permissible concentrations in hydrogen for automotive fuel cell use of 0.2 and 2 $\mu\text{mol mol}^{-1}$ of carbon monoxide and carbon dioxide, respectively.⁴⁰ Discussion in this section is restricted to the impact of organic impurities on the performance and stability of the catalyst layer; we are not aware of any studies on their impact on corrosion resistance of PTL and BPP components.

3.4.2 Adsorption and dissolution. Organics are well known to be oxidised at the anode of PEMWEs, producing carbon monoxide or carbon dioxide. The thermodynamic potential for these reactions to occur is around 0.1 V *vs.* RHE, much lower than that of the oxygen evolution reaction that normally operates at the anode. This suggests that these reactions should occur preferentially and lower the voltage required for an electrolyser to achieve the same current density. As such, there is a body of research focused on operating PEMWEs with hydrocarbon feeds to achieve lower energy electrolysis than is

possible with pure water. Readers are referred to reviews in the area.^{43,44,190} Methanol is the most common compound studied for this application, in concentrations typically at, or in excess of, 4 M with cells using either platinum or platinum–ruthenium catalysts on the anode and cathode. Measurements were typically performed in single cell and small stack configurations. The electrolysers achieved significantly lower voltages than water-fed PEMWEs, with 1.67 A cm^{-2} at 1.2 V using 4 M methanol at 80 °C (ref. 191) in a single cell. A 5 cell, 50 cm^2 stack operated on 4 M methanol at 80 °C and 1 A cm^{-2} required a mean cell voltage of only 0.79 V, without notable increase in cell voltage for 2500 h.¹⁹² There are few reports of cells operating on more complex hydrocarbons or alcohols, with the kinetics of the process less favourable due to the need for the C–C bond activation.¹⁹³

However, the kinetics of methanol oxidation are much less established on iridium or ruthenium oxide electrodes than for platinum containing electrodes. Typically, both ruthenium and iridium oxides require potentials >0.9 V *vs.* RHE to yield any measurable methanol oxidation activity.^{194,195} We anticipate that at typical PEMWE anode operating potentials (>1.5 V *vs.* RHE), there is sufficient driving force for the oxidation reaction to take place and trace impurities are unlikely to remain unoxidized. Indeed, certain types of water purification system use PEMWE-style devices to oxidise organics in wastewater.^{196,197} Using simulated contaminated water, Petrov *et al.* demonstrated that low concentrations of organic species are oxidised without significant effect on the performance of a PEMWE, utilising a $\text{Ru}_2\text{IrTaO}_x$ anode catalyst and platinum cathode catalyst.¹⁹⁶ The polarisation curve showed very little difference between pure water and pure water with 40 ppm TOC (a mixture of aromatic and non-aromatic organic alcohols and acids). In addition, the oxidation process was demonstrated to occur at high rates during OER with carbon dioxide detected using gas chromatography of the anode outlet gases. More challengingly, the formation of carbon monoxide and carbon dioxide *via* oxidation of organic molecules during operation introduces additional impurity into the PEMWE system. While this is likely not problematic at the anode, crossover to the cathode could prove detrimental as CO_{ads} can easily form and block active sites of the platinum catalyst at cathode.¹⁹⁸ At sufficiently low potentials (0.06 V_{RHE}), carbon dioxide may also be reduced to CO_{ads} , providing a further source of ECSA reduction.

Organic species containing certain functional groups, such as –OH and –CN, can form stable metal complexes with dissolved metal species.¹⁸⁹ This effect can be further enhanced by chelating agents such as ethylenediaminetetraacetic acid (EDTA), which can enhance dissolution of many metals, including PGMs.^{199,200} The formation of these metal complexes could enhance dissolution of the catalyst by stabilising the dissolved metal species and preventing redeposition of the metal, leading to increased loss of catalyst performance. Nonetheless, the extent to which this phenomenon affects performance at both the small scale experimental (*i.e.*, RDE) and the full system level (*i.e.*, single cell/stack) is unknown. A recent study conducted by Kormányos *et al.* on fuel oxidizing PtRu/C catalysts demonstrates the effect of organic species on



platinum and ruthenium dissolution rates, using online inductively coupled plasma mass spectrometry.²⁰¹ In 0.1 M HClO₄ the presence of different organic fuels (methanol, ethanol, and isopropyl alcohol) had no discernible effect on the onset potential for platinum or ruthenium dissolution. However, the presence of carbon monoxide did shift the dissolution potential to more negative values by 130–140 mV; we attribute this enhancement of dissolution in the presence of carbon monoxide to the roughening it can induce to precious metal catalysts,^{202,203} in turn exposing more under-coordinated surface sites, which are the most prone to dissolution.²⁰⁴ It is not clear either why this mechanism does not occur when operating on methanol as CO_{ads} is a key intermediate or how this phenomenon may affect iridium oxide or platinum electrocatalysts.

Both platinum and ruthenium are found to dissolve when cycled to potentials of 1.2 V vs. RHE or higher, and this is enhanced in the presence of methanol, as shown in Fig. 10.²⁰⁵ We anticipate that oxide catalysts, such as iridium–ruthenium oxides or even perhaps iridium oxides, may also be subject to enhanced dissolution in the presence of methanol under PEMWE conditions.

3.4.3 Inert impurities. Solid, inert, particles may be introduced into the water from external sources and/or failure of the water purification system. These particles can accumulate and block the PTL pores, leading to mass transport problems. Silicon particles have been shown to increase in concentration from the usage of polymer-based water tanks. One study showed

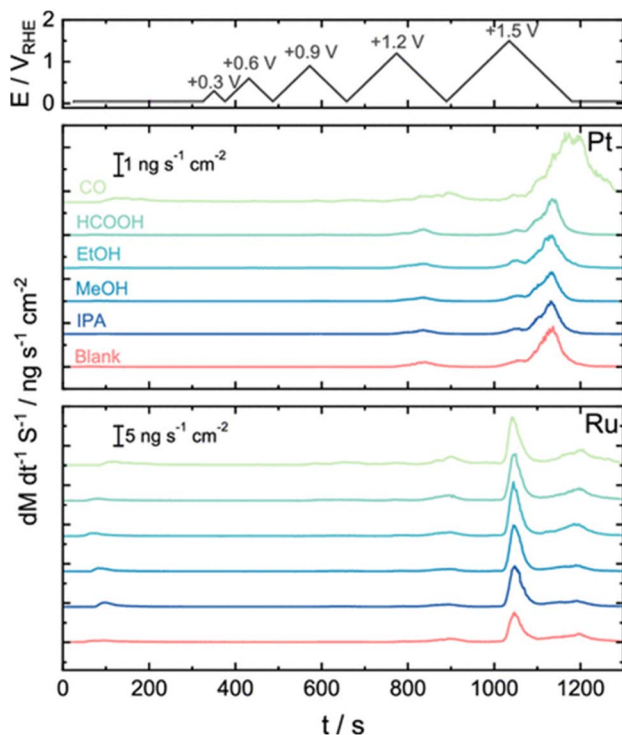


Fig. 10 Results from online ICP-MS using a scanning flow cell indicating the effect of 0.05 M organic fuel contamination on catalyst dissolution. This figure has been reproduced from ref. 201 under CC-BY-NC-ND licence.

that after operation of a PEMWE for 204 h, the circulated water showed an increase of silicon, sodium, calcium, NO₃⁻, and SO₄²⁻.²⁰⁶ Silicon likely originated from the nano-silica filled polypropylene tank, while SO₄²⁻ and NO₃⁻ were suspected to arise from oxidation of microbes at the anode, which was consistent with the observed decrease in TOC with time. Unfortunately, there is a lack of studies on the impact or likelihood of microbial impurities on PEMWE performance. At the anode, low quantities of microbes would likely be oxidised; it's not clear this is the case at the cathode, however. Studies on inert solid impurities are still lacking, and further experiments are required to verify their impact.

3.5 Summary and outstanding research challenges

3.5.1 Summary. There are four main types of water impurities identified for PEMWE: cation, anion, organic, and inert. The type of impurity, possible route of entry, and impact on PEMWE components are summarised in Table 2. Most cations fall under the category of exogenous impurity, while only titanium and PGM cations are endogenous. Cations degrade performance mainly through substituting the protons within the membrane and ionomer. At the catalyst layer, this leads to decreased proton pathways and slows down the electrochemical reaction. In the membrane, charge carriers are substituted with less mobile ions, thereby lowering the ionic conductivity. Iron cations can also trigger the Fenton reaction, initiating membrane degradation.

Anions are mainly sourced exogenously, except for F⁻ and SO₄²⁻, which are products of membrane degradation. Anions are less damaging in comparison to cations as they cannot substitute protons and lower ionic conductivity. Most anions can be adsorbed on the catalyst surface, resulting in reduced activity. Of all anions, chlorides are the most widely studied as they are present at higher concentrations in seawater and tap water. Moreover, chlorides can be oxidised into chlorine at the PEMWE anode. This can lead to secondary damage to tubing and ancillary equipment, and contamination of product gases. In principle, other anions can be adsorbed on the catalyst but the low potential at the cathode (<0 V vs. RHE) and high potential at the anode (>1.5 V vs. RHE) usually prevent this issue.

Organics and inert impurities are the remaining two of the possible exogenous impurities in the PEMWE water. 'Organic' is a very broad classification and different functional groups can induce unique effects. Groups such as -OH and -CN may induce chelating effects and promote PGM dissolution but, most commonly, organics adsorb at the catalyst surface and block active sites. At the anode, organics such as methanol can be expected to be oxidised, but gaseous oxidation products can cross over to the cathode, lowering the hydrogen quality and poisoning the cathode catalyst. For inert impurities there is little research performed to date and the most feasible issue is solid particulates introduced by the BoP, leading to mass transport issues.

3.5.2 Outstanding research challenges. Further work is required to understand the level of impurities likely to be



Table 2 Impact of impurities classified by their source and impact on PEMWE components

Impurity	Source	Cell components	Membrane	Anode	Cathode
Na ⁺ , Mg ²⁺ , Ca ²⁺	Seawater, potable water		Lowers membrane conductivity ^{95,96,104,106,107,131,159}	Lowers ionomer conductivity, ²⁰⁷ catalyst poison ⁴⁷	Lowers ionomer conductivity, ⁴⁷ decreases rate of HER ^{57,123,126,127,129}
Ti ⁴⁺ Ni ²⁺ , Cu ²⁺ Fe ²⁺ , Fe ³⁺	PTL corrosion External contamination Stainless steel tubing degradation		Lowers membrane conductivity, degrades membrane ^{106,107,111,124,125,207,208}		
Ir ²⁺ Cl ⁻	Anode catalyst dissolution Water purification failure			Enhances dissolution, ^{170,182} catalyst poison, unwanted side product ^{147,182,209} Enhances dissolution ¹⁸⁵⁻¹⁸⁸	
F ⁻	Water purification failure, membrane degradation	Corrodes Ti PTL and BPP ^{54,137}			
Br ⁻ , I ⁻	Water purification failure			Enhances dissolution, ^{155,156} catalyst poison ¹⁵⁷⁻¹⁵⁹ Catalyst poison ¹⁶⁶	
SO ₄ ²⁻	Water purification failure, membrane degradation			Enhances dissolution, ^{199,200,205} catalyst poison, ¹⁹⁸ unwanted side product ^{194,195}	
Organics	External contamination			Enhances dissolution, ^{199,200,205} catalyst poison, ¹⁹⁸ unwanted side product ^{194,195}	
Solid particulates	External contamination			Blocks PTL pores and induces mass transport issues ²¹⁰	

encountered in real PEMWE systems as even trace impurities may be important over long operating lifetimes. For instance, there is little information on the trace solubility of BoP materials in ultra-pure water and on the real composition of deionised feed water in commercial scale systems. There is also a scarcity of impurity studies performed under industrially relevant PEMWE conditions and cell hardware. The only studies at single cell level are for cations, primarily Na⁺ and Fe²⁺ with the chosen concentration often quite high (>1 mM) and unrepresentative of those likely to be encountered during real world operation. On the anion side, most studies have been performed in a three-electrode cell setup. This work needs to be extended to single cell level to assess whether trace CER accelerates degradation in PEMWEs. Chloride-related studies have typically been framed from the direct seawater electrolysis perspective and thus have tended to use a very high Cl⁻ concentration. In PEMWEs, Cl⁻ may originate from slow deterioration of the purification processes or accidental contamination, *e.g.*, from process upsets. Although the relevant concentration is still undefined for such incidents, concentrations within the ppm range might be expected. Further likely developments in PEMWEs need to consider the impact of trace impurities; for instance, the trend of moving towards more active catalysts with lower loadings and higher ECSA is likely to make them more susceptible to both poisoning and dissolution.

The impact of organic impurities is another topic that is barely touched upon in the literature. The primary industrial context for water-containing organics is the electrooxidation of wastewater, so the reported data are often focused more on the organic oxidation efficiency and less on cell performance. There

is a clear economic benefit in using lower quality water as it would simplify the associated cost of water purification or reduce the energy required for hydrogen production *via* electrolysis. The underlying premise is that organics should not interfere with PEMWE performance as they would simply be oxidised. However, some concerns remain, such as the potential for crossover carbon monoxide and carbon dioxide to lower the quality of hydrogen and adsorb on the cathode catalyst. More detailed studies would help to verify these concerns and could aid the development of organic fed water electrolysis technology. Finally, the impact of solid and microbial impurities on PEMWE performance has yet to be assessed. In the event of rupture of the reverse osmosis membrane, these solid inert impurities could enter the water and disrupt the stack performance, but simple mitigation or recovery approaches may be possible.

4 Impact of impurities in AWEs and AEMWEs

4.1 Introduction

This section addresses the impact of endogenous and exogenous impurities at both the anode and cathode of AWEs and AEMWEs. As in the PEMWE section, impurities are classified into three main types: cationic, anionic, and organic, with plausible mechanisms of operation and impact assessed for both technologies simultaneously, as summarised in Fig. 11. For many classes of impurity there is limited quantitative information available, so the discussion again remains largely qualitative.



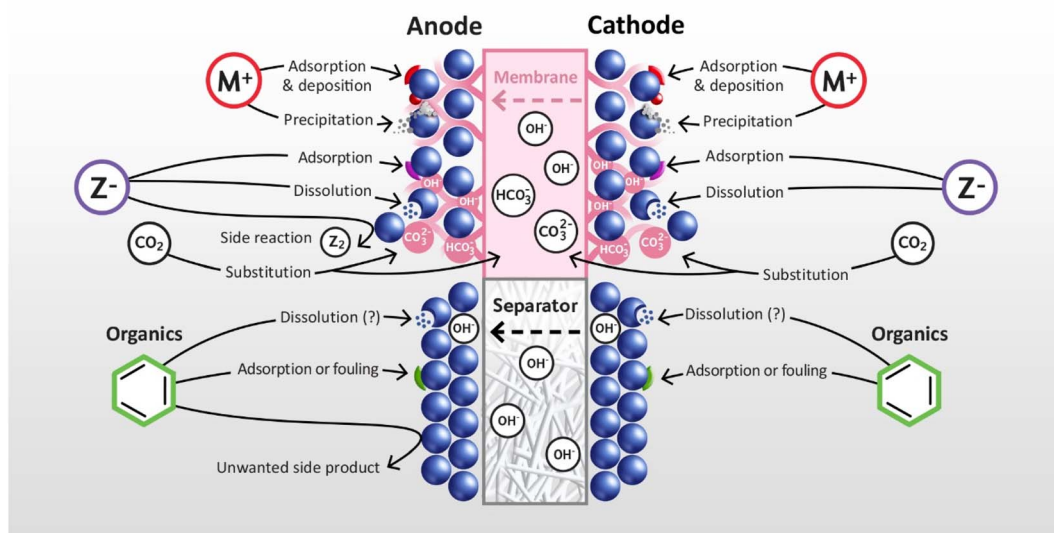


Fig. 11 Schematic summary of the impact of various types of impurity on AEMWEs (top) and AWEs (bottom).

4.2 Cationic impurities

The main charge carriers in AWEs and AEMWEs are anions, particularly OH^- . Therefore, cations do not pose a direct threat to the membrane, ionomer, and electrolyte as they do in PEMWEs. In fact, some of the cationic impurities that can commonly be found in water and potassium hydroxide electrolyte actually enhance the performance of some OER catalysts. The two main mechanisms by which cationic impurities can affect performance are adsorption on the catalyst and deposition in the solid phase.

4.2.1 Adsorption and deposition. Reagent grade potassium hydroxide is well known to contain ppm levels of impurities such as Ni^{2+} and Fe^{2+} . Corrigan and Bendert assessed the impact of iron impurities on the OER activity of a nickel oxide catalyst. After 10 charge/discharge cycles, the overpotential at 8 mA cm^{-2} in purified potassium hydroxide increased, while a decrease of overpotential was observed in potassium hydroxide with 1 ppm Fe added. Deposition of iron on the catalyst surface was the main reason for this improvement; just 0.01% iron was shown to decrease both OER overpotential and Tafel slope.²¹¹ Iron, cerium and cobalt co-precipitated on $\text{Ni}(\text{OH})_2$ at >10 wt% were also reported to increase OER performance while lead, zinc, and cadmium poison the OER by increasing the overpotential at 16 mA cm^{-2} .^{212,213}

One of the mechanisms by which deposited iron improves the performance of $\text{Ni}(\text{OH})_2/\text{NiOOH}$ electrocatalysts is by increasing the electrical conductivity by up to a factor of 30.²¹⁴ However, the main reason behind the increased OER activity is apparently changes to the active site. Without any iron on the $\text{Ni}(\text{OH})_2/\text{NiOOH}$ catalyst, $\gamma\text{-NiOOH}$ was reported to be the most active phase for OER.²¹⁴ Iron incorporation on $\text{Ni}(\text{OH})_2$ was observed to take place continuously in non-purified potassium hydroxide, with one experiment showing that the iron content increased to 23–26 at% of total metal content after 38 days of aging. The overpotential at 10 mA cm^{-2} was observed to reach

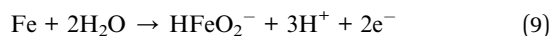
a minimum at 12–17 at% iron after 5 days of aging. It was found that the deposited iron formed a NiFe layered double hydroxide structure,⁸⁸ which is one of the most active non-precious metal OER catalysts in an alkaline environment reported to date.²¹⁵ Intentionally adding iron salt such as FeO_4^{2-} into the electrolyte was also reported to improve OER activity.²¹⁶ When cycled from -0.14 V to $1.86 \text{ V vs. Hg/HgO}$ using reagent-grade potassium hydroxide, an increase of OER current by a factor of 1.75 was observed after 150 cycles. Addition of K_2FeO_4 after the 150th cycle further increased the OER current by a factor of 1.1. Apart from iron, other metallic impurities can also play an active role in the performance of AWE systems. Fluorine-doped tin oxide on glass was used as an electrode to perform OER in a 0.5 M BK_3O_3 solution with metallic impurities. Initially no current was generated at 1.4 V vs. NHE , yet after prolonged anodization a rise of current was detected. The 17 nM concentration of nickel within the electrolyte was reported to be deposited and acted as an active catalyst. It was demonstrated that when amberlite resin was used to clean the electrolyte, the increase of current was smaller in magnitude.²¹⁷

At the cathode, conflicting reports have been published on the impact of iron deposition on the performance of nickel catalysts. When a nickel wire was held at $-1.5 \text{ V vs. Hg/HgO}$ in 30% KOH with 0.5 ppm iron for several hours, the overpotential was observed to increase continuously.²¹⁸ *Post mortem* analysis revealed iron deposition and thus one of the causes of degradation was attributed to iron being a worse catalyst for HER than nickel. However, measurement of a Ni200 cathode (99.6% purity) showed contradictory results. The Ni200 cathode was tested at 37°C , 250 mA cm^{-2} for 10^5 s in 30 wt% potassium hydroxide with the addition of either 0.03 ppm or 3 ppm iron. The cathode degraded at the same rate in both solutions until around 10^3 s , and the electrode in 3 ppm iron experienced a recovery to its original potential after 10^5 s . The electrode in 3 ppm iron was found to change colour to black and to have



picked up iron impurities. The performance increase was then attributed to an increased active surface area.²¹⁹

One of the possible reasons for the decrease in nickel performance during HER is hydride formation. In a specially designed permeation cell, nickel foil was secured between two compartments, with HER taking place in one compartment and the hydrogen diffusing through the metal to the other compartment, where a permeation current is recorded. In 30% potassium hydroxide, at charging currents of 25 mA and 50 mA, the hydrogen permeation current firstly increased and then dropped as the HER overpotential increased. This was attributed to the formation of beta nickel hydride on the surface, which acts as hydrogen diffusion barrier. The decrease in electroactivity was attributed to the d-band filling of nickel in beta nickel hydride.²²⁰ Nickel hydride was also observed on a RANEY® Ni–Al anode after operation at 300 mA cm⁻² in 10% potassium hydroxide. A continuous increase in overpotential was observed and X-ray diffraction (XRD) revealed hydride formation. Post-HER, the electrode was annealed under argon at 1000 °C for 2 h, which significantly reduced the overpotential and removed nickel hydride.²²¹ Iron deposition was reported to mitigate this hydride formation at the cathode. Nickel foil with 0.04 μm of sputter deposited iron on the surface was found to have a stable overpotential and stable hydrogen crossover current after 45 h in the permeation cell, in contrast to bare nickel.²²² It has been shown using the permeation cell experiments that iron particles detached from the electrode form HFeO₂⁻ by corrosion, which can be reduced back to iron on the electrode surface at more negative potentials, as shown in the following reaction.²²³



This freshly deposited iron was proposed to increase the HER activity of iron-coated nickel cathodes. For nickel–iron cathodes, HER activity was reported to increase with iron content peaking at 90 wt% iron. Cathodic sweep experiments showed signs of reduction of Ni²⁺, Fe²⁺ and/or NiFe₂O₄ to nickel and iron, which in turn enlarged the active surface area. However, corrosion did take place on all nickel, iron, and Ni–Fe samples forming oxide layers. The cathodic potential was proposed to reduce these oxides into active intermediates such as adsorbed FeOH and adsorbed NiOH that further enhance HER activity.²²⁴

A similar increase of HER performance after iron or nickel deposition has been reported for silver, gold, copper²²⁵ and cobalt based catalysts.²²⁶ Improvement in HER activity of copper and ruthenium based catalysts due to leaching and deposition of titanium oxides from a titanium current collector has also been reported.^{80,227} Nickel hydroxides either introduced as an impurity in the electrolyte or deliberately deposited have also been shown to improve the catalytic activity of nickel, silver, copper and platinum based catalysts.^{228,229} The oxide and hydroxide-based impurities discussed above have been shown to improve the catalytic activity by optimizing the water dissociation step of HER. It is clear that any metallic impurities that

are active for OER and HER reaction would likely not induce degradation upon deposition.

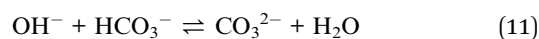
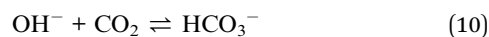
While some metallic impurities are beneficial for nickel-based cathodes, others can prove detrimental. Zn²⁺ is often present in low amounts in potassium hydroxide electrolyte and can also leach out from plastic components and gasket materials. Zn(NO₃)₂·6H₂O at 0.067 mg mL⁻¹, which is significantly higher than the trace ppm level found in potassium hydroxide, was added to the catholyte of a nickel/nickel electrolyser operated at 50 mA cm⁻². The cell voltage increased from 2.6 V and reached a stable 3.1 V for the next 6 h of operation. Clear zinc dendrite deposits were identified post operation. Zinc was reduced at -1.43 V vs. Ag/AgCl/3 M KCl, while the cathode HER potential at 50 mA cm⁻² was -1.55 V vs. Ag/AgCl/3 M KCl.²³⁰ Synthetic tap water was also reported to degrade the HER rate. The synthetic tap water consisted of mg L⁻¹ levels of the following impurities: Na⁺, Ca²⁺, Mg²⁺, Al³⁺, Cl⁻, HCO₃⁻, SO₄²⁻, and NO₃⁻. Held at 0.25 A cm⁻², the performance of the electrolyser degraded continuously during 55 h of operation, with the anode potential increasing from 1.8 V to 2.4 V vs. RHE. Interestingly, gas product analysis revealed no significant difference between using pure water and synthetic tap water. EDX of the nickel electrode surface showed the presence of carbon (39.82%), oxygen (15.39%), iron (28.73%), lead (4.78%), sodium (2.97%), chlorine (2.19%), potassium (3%), and zinc (1.7%).³⁸ Most of these metallic impurities were not included when creating the synthetic tap water; therefore zinc, lead, and iron must have originated from salt impurities. Reports have shown that Ni–Fe²¹⁵ and Ni–C²³¹ are active catalysts for HER, so the most likely culprits for the deactivation are lead, zinc, and Cl⁻.

4.3 Anionic impurities

In both AWEs and AEMWEs OH⁻ is usually the charge-carrying ion, though we note some AEMWEs use alternative or mixed anions. Potassium hydroxide at concentrations above 20 wt% is widely used as the circulated electrolyte in AWEs. In theory, AEMWEs may operate on pure water with the hydroxide in the anion exchange membrane transporting charge, however contemporary practice is to use a low concentration 0.1 M to 1 M supporting electrolyte to obtain higher performance and durability, with aqueous potassium hydroxide being the most common choice. The main mechanisms by which anionic impurities can impact AWE and AEMWE performance is through charge carrier substitution in the electrolyte and adsorption on the catalyst layer.

4.3.1 Substitution

4.3.1.1 Carbon dioxide. An impurity of concern in an alkaline environments is carbon dioxide. Carbon dioxide is present in ambient air at around 400 ppm and can readily react with potassium hydroxide to form bicarbonate (HCO₃⁻) and carbonate (CO₃²⁻) anions according to the following reactions:



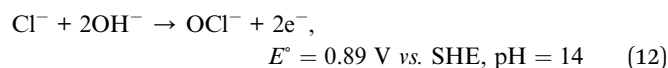
The radii of the hydrated forms of HCO_3^- and CO_3^{2-} are 3.64 Å and 3.94 Å, respectively.²³² The carbonates are larger in size than OH^- at 3 Å, which in turn lowers their overall ionic mobility. Relative to K^+ , the mobilities of OH^- , HCO_3^- , and CO_3^{2-} are 2.71, 0.6, and 0.98, respectively.²³³ The presence of carbon dioxide lowers the overall cell performance of an AWE by reducing the effective electrolyte conductivity. The impact of carbon dioxide on the performance has been studied extensively in alkaline fuel cells (AFCs) which are necessarily exposed to ambient air in a way that electrolyzers are not. A half-cell AFC study using 7.1 M potassium hydroxide and 5% carbon dioxide showed a linear increase of carbonate concentration during 3000 h of operation. Interestingly, this increased concentration of carbonate did not affect the performance of the electrode, as evidenced by similar performance degradation of control (pure H_2 fed) and 5% CO_2 containing H_2 fed electrodes. The decrease of cell performance was mainly attributed to the decrease of electrochemical surface area of the catalyst over time.²³⁴ In another study using a microfluidic fuel cell, a critical ratio of hydroxide/carbonate to cause a 10% voltage decrease was proposed.²³⁵ In the case of 3 M potassium hydroxide, the critical potassium carbonate concentration was 0.75 M (reached after 1700 h operation) while for 5 M and 7 M potassium hydroxide it was 1.75 M (reached after 3900 h of operation). The absence of significant conductivity decline in ref. 234 was simply because the critical concentration of carbonate was not reached. Carbon dioxide contamination in AWEs should be much slower in sealed systems with water purification, so timescales for contamination should be longer.

In the case of AEMWEs, there have been various studies on anion exchange membrane fuel cells (AEMFCs) using similar AEMs.^{236–238} For three types of AEM with different anion forms (OH^- , HCO_3^- , CO_3^{2-}), the conductivity was found to decrease in the following order: $\text{OH}^- > \text{HCO}_3^- > \text{CO}_3^{2-}$ with increased relative humidity and membrane hydration reported to remedy the conductivity decline.²³⁹ It is likely that the extent of carbon dioxide poisoning in AEMWEs, where water fully hydrates the membrane, is not as severe as that reported in AEMFCs; additionally, OH^- is generated *in situ* by simply running alkaline water electrolysis reactions and this will gradually purge the carbonate as carbon dioxide.²⁴⁰ This has been demonstrated in AEMFCs when deliberately passing a low electrolysis current or operating at high current densities.²⁴¹ In case of OER, the presence of carbonate was found to decrease the catalytic activity of LaNiO_3 by formation of carbonates which altered the chemical environment of active sites.²⁴² This may be generally applicable to other transition metal based OER catalysts.

Apart from decreasing the ionic conductivity and catalyst activity, carbon dioxide has been reported to affect membrane physicochemical properties such as membrane crystallinity and side-chain spacing.²⁴³ The reaction to form CO_3^{2-} generates water, which then leaves the membrane; this shrinks the side-chain spacing and affects membrane crystallinity. However, a higher humidity (RH: 85%) was reported to reduce the extent of this polymer morphology change. Carbon dioxide contamination of the water tank was also a problem for a pure water operated AEMWE.²⁴⁴ Polarisation curves of the cell in quick

succession showed rapid degradation while the HFR increased 3-fold from $0.6 \Omega \text{ cm}^2$ to $1.67 \Omega \text{ cm}^2$. This short-term degradation was not observed when measures to minimise carbon dioxide ingress, *i.e.*, placing the whole setup inside a glovebox, were put in place, clearly illustrating the importance of effective gas-tight seals when operating AEMWEs with pure water.

4.3.2 Reactions. In contrast to acidic electrolysis, chloride anions are less problematic for alkaline operation. There is a much larger offset (480 mV) between the CER and OER at alkaline pH in comparison to that under acidic conditions (*ca.* 200 mV). As the Pourbaix diagram in Fig. 8 shows, the thermodynamically stable product of chloride oxidation shifts with pH. When the pH is higher than 8, the following reaction takes place:



The 480 mV overpotential window before the initiation of reaction (12) has routinely been exploited to facilitate operation of direct seawater electrolysis without generation of chlorine or OCl^- at the anode.³⁷ It is not within the scope of this article to explore in-depth the issue of seawater electrolysis; the reader is referred to other excellent reviews of the subject.^{37,142} However, most studies regarding the issue of chloride contamination have been in the context of seawater electrolysis applications.

NiFe layered double hydroxide (LDH) catalyst, a common AWE OER catalyst, was reported to be able to perform OER close to 100% faradaic efficiency in 0.5 M sodium chloride + 0.1 M potassium hydroxide.¹⁴⁸ Keeping the pH at 13 with an overpotential of 480 mV, 10 mA cm^{-2} was achieved without any chlorine production. A 2 h stability test at 10 mA cm^{-2} under different pH showed degradation of NiFe LDH at pH 9.2 with and without sodium chloride. A sharp increase of potential from 1.7 V to 2.4 V vs. RHE was observed after 60 min of operation with sodium chloride. When both sodium chloride and potassium hydroxide were used, the cell potential remained below the 480 mV limit during 120 min of operation. The same catalyst was also proven to work well in a AEMWE configuration with a Pt/C cathode and Tokuyama A201 membrane.²⁴⁵ Three concentrations of potassium hydroxide (0.1 M, 0.5 M, and 1 M) were tested with and without addition of 0.5 M sodium chloride by setting the cell voltage at 1.6 V. All cells showed degradation over a 100 h period, except that in 0.1 M potassium hydroxide without sodium chloride. Scanning electron microscopy and impedance spectroscopy indicated no degradation of the electrode or membrane thickness with or without sodium chloride. The evidence suggested that the degradation was mainly due to chloride ions substituting hydroxyl ions and lowering conductivity. Interestingly, the assumption of a 480 mV window of operation is only valid from a thermodynamic perspective. Using a symmetrical feed of 0.5 M potassium hydroxide and sodium chloride at both cathode and anode, it took a cell voltage $>2.4 \text{ V}$ to initiate any OCl^- formation using Ir-black and Ir/TiO₂ as the anode catalyst. When NiFe LDH was used as the anode, prolonged operation at high current densities ($0.2\text{--}0.7 \text{ A cm}^{-2}$) corresponding to cell voltage of $2.0\text{--}2.4 \text{ V}$ yielded a stable

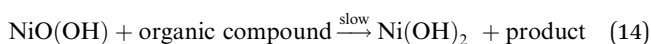
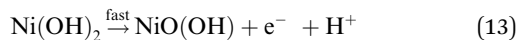


94% faradaic efficiency, suggesting only a minor contribution from the CER. Iodometric analysis of the electrolyte showed no OCl^- formation, demonstrating selectivity of NiFe LDH in the presence of Cl^- .²⁴⁶

4.3.3 Corrosion. Even without the CER taking place, nickel electrodes are susceptible to degradation in saline water operation.²⁴⁷ A nickel electrode (Ni200) operated in saturated sodium chloride + 30 wt% sodium hydroxide at 467 mA cm^{-2} failed after 9 days of operation, due to short-circuiting of the electrode from black particle formation. Upon inspection, the black particles were found to be NiO, a product of nickel corrosion in the highly saline electrolyte. Cl^- was deduced to attack the passive layer formed on the surface of Ni200. Certain configurations of nickel such as $\text{NiS}_x/\text{NiFe-Ni}$ foam anodes were reported to have better tolerance to operation in saline electrolyte. Four types of nickel catalyst (nickel foam, NiFe-Ni Foam, NiS_x -Ni foam, and NiFe/ NiS_x -Ni foam) were tested at 400 mA cm^{-2} , 1 M potassium hydroxide + 2 M sodium hydroxide (four times the chloride concentration of seawater concentration as an accelerated stress test).²⁴⁸ Nickel foam and NiS_x performance dropped almost immediately, followed by NiFe (12 h) and NiFe/ NiS_x (600 h). In another milder test (1 M potassium hydroxide and real seawater), NiFe/ NiS_x was reported to be stable for 1000 h of operation. XCT scans post operation showed that the source of degradation was the nickel being corroded. NiFe/ NiS_x paired with Ni-NiO-Cr₂O₃ at the cathode operating at $400\text{--}1000 \text{ mA cm}^{-2}$ showed 100% faradaic efficiency, agreeing with ref. 246 that the 480 mV operation window is not absolute.

4.4 Organic & inert impurities

4.4.1 Organic impurities. The oxidation of organics in alkaline systems has been investigated for AFCs, and several reports have been published on the ability of nickel catalysts to oxidise organic fuels such as alcohols. In the case of methanol, the structure of the catalyst was reported to play an important role for electrooxidation of organics. Smooth nickel, for instance, was reported to be very inactive due to the presence of superficial oxides, as compared to nickel-zinc alloys where the superficial oxides are easy to reduce.²⁴⁹ Nickel on carbon (Ni/C), however, was reported to initiate the methanol oxidation reaction in 1 M potassium hydroxide at 1.25 V vs. Hg/HgO/1 M KOH (MMO), a higher potential than that for initiation of the OER.²⁵⁰ Fleischmann *et al.*²⁵¹ proposed that the mechanism of oxidation of organics in nickel electrodes follows these reactions:



Various papers have pointed out that NiO(OH) is the required species for oxidation of organics, based on the fact that the oxidation potential coincides with the potential for NiO(OH) formation.^{252,253} One experiment showed that after cycling the potential (-0.9 V to 1.6 V vs. MMO) of Ni/C for 50 cycles, the magnitude of the methanol oxidation peak decreased by 65%, which was suggested to be as a result of nickel oxide formation.

In commercial AWEs and AEMWEs with nickel-based catalysts, even though the potential of the anode would be high enough to initiate oxidation, it seems that catalyst passivation may prevent complete oxidation of the organics.

Another issue is the nature of the by-products of these organic fuels upon oxidation. One study of a Ni[salen] complex catalyst at different applied potentials showed that the oxidation reaction starts at 0.4 V vs. Ag/AgCl and generates a range of products. Methanol and glycerol are mainly oxidised into formate, while ethanol held at 0.6 V vs. Ag/AgCl forms acetate instead.²⁵⁴ In AWEs and AEMWEs, the anode potential will be much higher than this, in which case oxidation of the organics into carbon monoxide and carbon dioxide can be expected. Carbon dioxide poses significant challenges in alkaline electrolytes as discussed above. As with PEMWEs, both carbon dioxide and carbon monoxide may also contaminate the product hydrogen.

The right type of organic could also impose a chelating effect on the nickel catalysts. For instance, EDTA and [S,S]-ethylenediamine disuccinic acid ([S,S]-EDDS) have been used as chelating agents in metal recovery.²⁵⁵ Their ability to form stable metal complexes may cause or accelerate catalyst degradation. This should be given careful consideration should wastewater containing unknown organics be used as the water feed. For alkaline systems, the presence of isopropanol, methanol, ethanol and formic acid actually resulted in a stabilisation effect as it was noted that even up to the upper potential limit tested, 1.5 V vs. RHE, acetone was the primary product of oxidation and not carbon dioxide.²⁰¹ The acetone is shown to stabilize the catalyst by irreversibly adsorbing on the surface, which prevents the dissolution of surface oxide layers during the reverse scan. The conclusion from this work suggests that the mechanism by which enhancement or stabilisation of catalysts occurs, *via* organic species, is very dependent on the specific surface adsorption and resulting oxidation pathway of these species. This indicates that studies on materials specifically relevant to OER in both acid and alkaline environments are necessary to determine whether organic species are having a significant effect on catalyst stability.

4.4.2 Inert impurities. The highly alkaline pH of AWEs and AEMWEs can cause precipitation of metallic ions due to the low solubility of their oxides³¹ and carbonates. For example, high concentrations of carbonate ions may even cause potassium carbonate to precipitate from the electrolyte (potassium carbonate solubility 110.5 g/100 g water at 20 °C).^{256,257} The main concern with precipitation is clogging of porous layers, such as the separator or electrodes, causing mass transport issues. However, such an issue would only be expected if high concentrations of carbonate or metal ions with low solubility were introduced, for example through failure of the water purification unit.

4.5 Summary and outstanding research challenges

4.5.1 Summary. AWEs and AEMWEs have three main types of impurities: cations, anions, and organics (Table 3). Cationic impurities may be sourced from the supplied electrolyte, the



water source or components leaching out. Most transition metal cations such as Fe^{3+} or Ni^{2+} actually have a beneficial effect when adsorbed on the catalyst, improving catalytic activity for both OER and HER. This is because these transition metals are active catalysts for HER and OER under alkaline conditions. Therefore, there is no urgent need to operate AWEs/AEMWEs using purified potassium hydroxide. However, there are still some cations, such as Zn^{2+} , Cd^{2+} , or Pb^{2+} , that are not active OER or HER catalysts and would decrease the performance upon adsorption. Ca^{2+} and Mg^{2+} cations can also deposit at high pH and may block reactant access to catalyst sites.

Since the charge carriers in AWEs and AEMWEs are anions, anionic impurities pose a significant threat. Carbon dioxide from ambient air can easily diffuse into the cell if there is a gasketing problem or a slightly loose connection within the stack. It forms HCO_3^- and CO_3^{2-} anions that lower the cell ionic conductivity and performance considerably. Moreover, carbon dioxide has also been reported to change the membrane physical properties in AEMFCs. Halogen ions can also be introduced from impure feed water or malfunction of the water purification unit. Cl^- , in particular, can trigger unwanted side reactions and generate OCl^- . If the electrode structure is incompatible, it could also trigger corrosion of the electrode.

There have been many studies of organic impurities in three-electrode setups, but all of them are framed from an alkaline fuel cell or organic electrooxidation perspective. Organic impurities can be introduced when lower quality water is used as the water feed. It is expected that at the anode, the organics will be oxidised into various products depending on the applied

potential. Side products such as formates and acetates have been reported. Other oxidation products, such as carbon dioxide and carbon monoxide, are likely but have not been explicitly reported. Gaseous and volatile products can contaminate the product hydrogen quality, leading to a requirement for additional purification processes.

4.5.2 Outstanding research challenges. Impurity studies in the alkaline area have tended to be entirely overlooked, even though some of the performance of AWEs can be credited to metallic impurities within the potassium hydroxide electrolyte. Some studies have addressed this issue by purifying the potassium hydroxide in order to help benchmark catalytic assessment more accurately. Following a similar logic, a study on the impact of bulk impure potassium hydroxide and purified potassium hydroxide in a single cell would also be beneficial to reveal whether impurities do impact durability positively. Intentional addition of Fe^{3+} or Ni^{2+} into the electrolyte should also be tested for impact on performance at single cell or stack level to further elucidate the role of these cations. Identifying other possible cations that may leach into the electrolyte from the BoP is also critical, as some cations can deactivate the catalyst layer upon adsorption.

On the anion side, optimisation of electrodes can lead to OER selective anode catalysts in the presence of chloride anions. This would be beneficial for direct seawater electrolysis but could also prove important if lower quality water were used to cut purification costs. A detailed study of the impact of carbon dioxide is also lacking for alkaline electrolyzers, as most papers report results within the context of fuel cells. Fuel cells

Table 3 Impact of impurities on AWEs and AEMWEs

Impurity type	Source	Anode	Separator/membrane/conductivity	Cathode
Fe^{3+}	Electrolyte impurity	Adsorbs on catalyst improvement of OER activity ^{88,211,214,216}		Improvement of HER activity ^{222–224} increased surface area ²¹⁹
Ni^{2+}	Electrolyte impurity	Adsorbs on catalyst. Improvement of OER activity ²¹⁷		Improvement of HER activity ^{228,229}
Al^{3+}	Electrolyte impurity	Adsorbs on catalyst. No adverse effect reported		No adverse effect reported
Zn^{2+}	Gasket leaching out, electrolyte impurity	Catalyst poison ²¹²		Forms dendrites at negative potentials, catalyst poison ²³⁰
Cd^{2+} , Pb^{2+} $\text{Ti}^{3+}/\text{Ti}^{4+}$	External contamination Electrode leaching out	Catalyst poison ²¹²		Improvement of HER activity ^{80,227}
Na^+	Water impurity (if NaOH is not used)	No adverse effect reported		No adverse effect reported
Ca^{2+} , Mg^{2+} Cl^-	Water impurity Water impurity	Deposits $\text{Mg}(\text{OH})_2$ at high pH, ^{31,258} may cause mass transport issues at electrodes Nickel corrosion, apart from special structures. ^{247,248}		
CO_2	Air contamination		Lowers conductivity, ^{232,234,235,237–239,259} reduces side chain spacing and membrane crystallinity ²⁴³	Increases charge transfer resistance (AEMWE) ²⁵⁹
Organics Inert impurities	Water impurity Precipitates formed by reaction with electrolyte	Catalyst poison ²⁵⁵	Clogging of pores ^{256,257}	Catalyst poison ²⁵⁵



operate with air and have a higher concentration of carbon dioxide, while electrolyzers would likely have a lower concentration. The concentration should be addressed in a more relevant range.

Finally, there is no organic impurity related study within the context of AWEs and AEMWEs. Many questions remain to be answered, including how electrooxidation could impact the performance of AWEs and AEMWEs and impacts on catalyst durability. Carbon dioxide, in principle, could be formed at high anode potentials and would lower the ionic conductivity of the electrolyte. There is still a steep learning curve to study whether organics could interfere with AWE and AEMWE operation in the long term.

5 Characterisation techniques, water purification technologies, and mitigation strategies

5.1 Characterisation techniques

A wide variety of analytical techniques are used in the testing of impurities and the effects on the materials involved for PEMWE, AEMWE and AWE. This testing can be split between half-cell investigations of small quantities of catalyst and single cell testing of complete systems. In addition to this, testing is split further between testing that can be done *in situ* (in a relevant environment), *in operando* (in an operating device) and *ex situ* (not in a relevant environment) with half-cell or full cell design being the limiting factor in the accessibility toward many currently available *in situ* and *in operando* characterisation techniques. This section provides a summary of the more commonly used techniques that could be used for relevant studies in the future.

5.1.1 Electrochemical techniques. Electrochemical testing is often carried out to determine the functional properties of electrolyzers or electrolyzer components, usually with a focus on determining performance or stability. This broad category may include both testing of devices as well as detailed testing of individual components.

Rotating Disc Electrode (RDE) or half cell testing are the most basic types of *in situ* electrochemical tests. Their main use is to characterise the behaviour of materials such as benchmarking the activity and stability of catalysts or other cell components. In a typical experiment the material under investigation is immersed in a liquid electrolyte along with a counter electrode and a reference electrode. A potential is applied to the material and the resulting current is measured. The benefit of these experiments is the well-defined potential of the sample, and ability to work with a range of electrode types including thin films of very small amounts of catalysts. A major limitation of these types of experiments is mass transport to and from the electrodes. In RDE, the electrode is rotated to precisely control the diffusion layer at the electrode surface but this is not often sufficient; for example the OER may form microbubbles that prohibit high current density operation.^{260,261} New cell designs, such as the floating electrode, that facilitate gas evacuation²⁶² alleviate mass transport effects. Half-cell testing often facilitates

the use of other *in situ* characterisation tools and it is easy to introduce impurities to the liquid electrolyte.

Rotating Ring Disk Electrode (RRDE) allows for additional *in situ* electrochemical probing of the process and in particular the intermediates that are formed at the working electrode.¹⁵⁰ While the material being investigated on the RDE is under current or potential control at the centre of a rotator an additional ring separated from the working electrode allows for a separate reaction to be probed by holding the ring at a different potential. This technique has been used to probe the quantity of chlorine being formed at an electrode, as well as demonstrating that oxygen evolution on iridium oxide is not suppressed by increasing Cl^- concentration.¹⁸² The chloride reduction reaction and oxygen reduction reaction can be competing reactions at the ring, so it was found to be necessary to select a condition that allowed only chlorine reduction: 0.95 V vs. RHE at pH 0.88.¹⁴⁷ The limitation of the RRDE technique is that it cannot distinguish the chemical identity of species at the ring electrode. Other analytical techniques are required to specifically identify the species and at what potential they are selective to certain reactions.

Single cell/full cell testing is carried out by operating a scaled down electrolyzer typically monitoring the cell voltage under a range of conditions. A single cell from a stack may be used or more commonly a small ~ 10 's cm^2 cell is assembled as a scaled down device.^{49,124} As all of the electrochemical components in an electrolyzer are present and acting as they do in a real device, the results are highly representative of real systems. However, this means that deconvoluting the response to determine mechanisms is challenging and usually relies on complementary techniques such as carbon monoxide stripping to determine ECSA, linear sweep voltammetry to determine hydrogen crossover, and electrochemical impedance spectroscopy to determine cell resistance and kinetic parameters.

Stack/system testing is the most relevant type of electrochemical testing, as this uses a real device, or one identically configured. Testing at this scale is resource intensive, time consuming and as with single cell testing it is challenging to deconvolute underlying processes. However, it can highlight phenomena that are only present in real systems, such as impurity enrichment, BoP degradation or stack design choices that exacerbate the impact of impurities.⁴⁷

Electrochemical Quartz Crystal Microbalance (EQCM) uses the high mass sensitivity of the resonant frequency of thin quartz crystal in an electrical field to measure changes in mass of a working electrode.²⁶³ These features allow EQCM to probe mass changes during *in situ* electrochemical half-cell experiments. This has allowed the investigation of the effects of impure potassium hydroxide on NiOOH films where iron contamination results in increased activity, but in particular the technique facilitated distinguishing between the effects of structure changes vs. iron incorporation that caused an increased mass of the electrode.⁸⁸ A limitation of this technique is that it cannot distinguish between different sources of mass loss such as dissolution and particle detachment.²⁶⁴

5.1.2 Material characterisation techniques. Due to the structure of the CCM many types of *in situ* characterisation can



be difficult to conduct so the major method for identifying both modes of degradation and contamination is through examination of the CCM both before and after testing. *Post mortem* analysis allows a range of techniques to be used to determine how the structure and composition change during operation.

Attenuated Total Reflectance – Fourier Transform Infrared Spectroscopy (ATR-FTIR) has been used to investigate degradation of the Nafion membrane in half cell tests.²⁶⁵ In particular, the ability of ATR-FTIR to investigate changing states of the functional groups within the membrane provides additional information about the impact of contaminants on membrane degradation that complements elemental mapping of contaminants. ATR-FTIR can also be used in half-cell tests on organic impurities and to reveal the by-products of oxidation of organics.²⁵⁴

Ultraviolet-visible spectrophotometry (UV-Vis) has been used for the quantification of the valence state of iron in cathode water by comparison of the Fe^{2+} signal recorded at 510 nm and using a standard concentration curve of prepared solutions to assess concentration.¹²⁴ In addition, *in situ* UV-Vis can be useful for detecting reaction intermediates in oxides²⁶⁶ as well as establishing the effect of various contaminants on surface species.²⁶⁷

X-ray absorption spectroscopy (XAS): X-ray absorption near edge spectroscopy (XANES), and extended X-ray absorption fine structure (EXAFS) typically use a synchrotron to provide a X-ray beam to excite core electrons.²⁶⁸ These techniques have been used to investigate the *in situ* effects of anion absorption of Cl^- , Br^- , OH^- , and HSO_4^- on platinum surfaces.¹⁵¹ While unlikely to be a commonly used technique for contaminant investigation in either PEMWE or AWE setups due to the requirement for a synchrotron facility as well as further limitations with cell design, it has proved insightful for fundamental studies of absorbed anionic species as well as demonstrating the capability of conducting *in situ* measurements.

Scanning Electron Microscopy (SEM)/Energy Dispersive X-ray Spectroscopy (EDX or EDS) involves coupling the detailed imaging of SEM with elemental mapping *via* EDS and is one of the common techniques used for *post mortem* analysis of CCMs. SEM provides different imaging modes with standard instruments usually offering both composition and topographical contrast and resolution on the sub μm scale. EDX has been used to particular effect both for the detection of cation contaminants and for analysing the amount of catalyst dissolved and where it deposits on the MEA.⁶⁸ Typically EDX has a detection range of 0.1–100 wt% of the material,⁶⁸ though the quantification of lighter elements such as oxygen and carbon is often unreliable, and the technique is not surface sensitive; as such it is better at detecting larger quantities of material deposited.

X-ray photoelectron spectroscopy (XPS) utilises the photoelectric effect, which involves firing X-rays at a sample and measuring the energy of emitted electrons, with each element giving a specific fingerprint signature.²⁶⁹ While *in situ* ambient pressure XPS has been used for the investigation of oxygen evolution materials^{267,270,271} it is more commonly used for *ex situ* characterisation of catalyst materials, where it is complementary to EDX with a high degree of sensitivity to the surface of

materials and quantification of lighter elements is more precise.

Electron probe microanalysis (EPMA), like EDS, utilises an electron beam and recording of emitted X-rays.²⁷² This provides spatial resolution on the sub-micrometer scale as well as the ability to detect concentration down to at least 100 ppm for most elements. This facilitates investigation of cross sections of CCMs for various cation contaminants such as Na^+ ,¹²² and Fe^{3+} .¹²⁴

Online neutron and X-ray imaging techniques utilise either a neutron or X-ray beam directed towards the sample, and the subsequent absorption and scattering of the radiation. The technique has often been employed to elucidate water-bubble dynamics within the flow field of PEMWEs.^{273–276} Applying neutron scattering to impurity studies is not straightforward, as it requires the impurity to have a sufficiently contrasting neutron scattering cross-section to water and other components. Certain ions, such as Gd^{3+} , fulfil this criteria by having an extremely high neutron scattering cross-section per atom, making them discernible.¹²³

Nanoscale Secondary Ion Mass Spectrometry (Nano SIMS) is an imaging technique that analyses the atomic mass of species on a solid sample surface with a resolution of 50 nm and a detection range in the sub-ppm range. The technique works by sputtering the top monolayer sample *via* a primary ion beam and analysing the ionised atomic and molecular species with a mass spectrometer. Nano SIMS has been routinely used for biological applications, but to date it has not been employed for *post mortem* CCM analysis.²⁷⁷ This technique offers a better resolution and detection for trace impurities and could be exploited in future research.

5.1.3 Chemical analysis of liquid phase impurities. Analysis of feed and recirculated water for contaminants is key to identifying both the concentrations of impurities and how these change over the course of electrolyser operation. In addition, it is useful for looking at degradation of materials *via* flow cell setups where the dissolved catalyst material can be accessed.

Inductively Coupled Plasma Mass Spectrometry (ICP-MS) is a type of mass spectrometry that uses inductively coupled plasma to ionise samples, allowing trace element analysis of liquid solutions.²⁷⁸ In the case of PEMWE and AWE research this can allow for analysis of recirculated water¹²⁵ or electrolyte for a range of contaminants. In addition, the technique can be used to monitor catalyst losses due to dissolution by analysing the quantity of dissolved catalyst in water/electrolyte in cell testing.²⁰¹ This may be of limited use for MEA analysis of flow water for anode catalyst dissolution as a majority of the dissolved anode catalyst will travel and deposit either in the membrane or on the cathode.⁶⁸ The detection limit of most elements in ICP-MS is also much lower than that of inductively coupled plasma optical emission spectroscopy (ICP-OES) with a lower limit of parts per trillion (ppt) for some elements/machines.

Scanning Flow Cell Inductively Coupled Mass Spectrometry (SFC ICP-MS) couples an electrochemical flow cell and ICP-MS allowing the composition of the electrolyte to be determined as a function of electrochemical properties. It allows *in situ*



studies of both acid and alkali cell setups, good time resolution and potential dependent studies.²⁷⁹ The high potential resolution enables the study of catalyst dissolution during potential scanning, allowing clear identification of the sensitivity to potential as well as the effect of contaminants on the quantity of catalyst that dissolves at a given potential.

pH monitoring encompasses a wide variety of techniques that measure pH. The most common methods are electrochemical, monitoring the potential difference between two electrodes.²⁸⁰ It is routinely used to monitor the quality of the feed water and acidity/alkalinity of prepared electrolyte. Monitoring the pH of the anode and cathode water in an electrolyser has been used to investigate the mechanism operating when poisoning with Na⁺ occurs.¹²¹

Ion chromatography measures the concentration of ions by separating them based on interaction with a resin and eluent. It is a technique that is often used to characterise ionic products in the water phase, particularly from membrane degradation, and can reliably detect impurities to the ppb level.^{141,281} While often used *ex situ*, incorporation of the ion chromatograph *in situ* to record the measurement of F⁻ ions during PEMWE operation has also been reported.²⁸

Polarography is an electrochemical technique that can be used to measure the concentration of metal ions in an electrolyte and can be incorporated in-line with continuous logging of data. The working electrode is either a dropping mercury cathode²⁸² or a hanging mercury drop electrode. Noble metal ions, such as platinum, can be measured in the ng L⁻¹ range²⁸³ and it has recently been used to measure the dissolution of PEMWE cathode material at open circuit with a limit of detection of 2 ng L⁻¹.⁶⁶

Ion selective electrodes are an off-the-shelf solution for measuring the concentration of specific ions. They work in a similar way to pH electrodes; instead of a proton selective membrane, a specific ion selective membrane is used.^{284,285} For example, a F⁻ ion sensitive electrode uses a LaF₃ crystalline membrane that only allows F⁻ to pass.²⁸⁶ These electrodes have been routinely employed for *in situ* and *ex situ* measurement of ions in PEMFCs, mainly for F⁻.⁸⁷ Their limit of detection is at the ppm level, so ICP-MS is preferred when reliable measurement at the ppb level is needed. However, such electrodes have the advantages of being cheap, robust, and modular.

Water conductivity is typically measured as a resistance between two electrodes placed in a water container of accurately known dimensions.²⁸⁷ In very pure water the self-ionisation of water contributes a significant proportion of conducting ions. Temperature influences the viscosity and ion movement which in turn affects the measured conductivity.²⁸⁸ Conductivity meters are routinely employed for on-line conductivity measurements of feed water in PEMWEs.²⁸⁹

Total Organic Carbon (TOC) is the amount of carbon present in organic compounds and is generally used as a non-specific indicator of water quality.²⁹⁰ TOC analysers measure amount of carbon dioxide formed by catalytic oxidation of organic content. The carbon dioxide is measured either using a conductivity cell (if it's in the aqueous phase) or by a non-

dispersive infrared cell. TOC measurements have recently been employed to measure MEA degradation in PEMWEs.²⁸⁹

5.1.4 Chemical analysis of gas phase impurities. In order to assess the purity of evolved hydrogen, techniques to measure contaminants in the gas phase can be used. These may also provide quantification of impurities; for instance, evolved carbon dioxide may be used as a means to identify oxidation of organic contaminants.

Gas chromatography (GC) can be used to determine the quantity of evolved gas *via* the use of an analytical column where different gases are separated by their interaction with the stationary phase.²⁹¹ This is particularly relevant for the study of both carbon dioxide and carbon monoxide as contaminant gases. The monitoring of these gases also allows for studying the effects of organic contaminant consumption at high potential operation where they are expected to undergo oxidation.¹⁹⁶

Differential Electrochemical Mass Spectrometry (DEMS)²⁹² allows for online detection of electrochemical products and intermediate species *via* a catalyst layer deposited onto a porous Teflon membrane supported on a glass or steel frit whose pores then lead to a mass spectrometer. The hydrophobic membrane prevents liquid from entering the pores but allows gaseous and volatile products to pass through. In addition, it can be used to semi-quantitatively determine the quantity of species produced. This technique has been applied to the study of Cl⁻ contamination on carbon supported iridium oxides to determine at the ratio of evolved chlorine to oxygen in conjunction with RRDE experiments.¹⁴⁷

On Chip Electrochemical Mass Spectrometry (EC-MS) is very similar to DEMS.²⁹³ This variant utilises a type of membrane chip that avoids the need for differential pumping, allowing faster time resolution and complete sampling of gaseous and volatile products/intermediates, providing a fully quantitative analysis. While it has not been used to date for any contaminant studies of OER catalysts, the orders of magnitude enhancement in sensitivity to molecules such as carbon dioxide and oxygen could make it highly useful to probe organic oxidation reactions.

5.2 Water purification technologies

Despite the critical role of water purity in the longevity of various low temperature electrolyser technologies, little has been openly published on the development of specialist water purification systems or techniques to recover performance loss due to poisoning incidents. The water purification requirements for PEMWE systems vary with manufacturer and installation, but in general most manufacturers stipulate a minimum feed water resistivity of 1 MΩ cm. AWE water purity is not often discussed in the literature, but a similar minimum value of 1 MΩ cm is to be expected as impurities such as Ca²⁺, Mg²⁺, and Cl⁻ need to be removed.

5.2.1 Reverse osmosis. The technology that is often employed for the first purification step is reverse osmosis (RO).²⁹⁴ If the water source is not tap water, *e.g.*, seawater or other sources of water, additional pre-treatment such as



sediment filters and activated carbon adsorption would also be installed before the RO membrane. A RO system works by applying external pressure to salt water over a semi-permeable membrane, allowing only water and trace amounts of mono-valent ions to pass through. Commercial RO membranes are predominantly organic polymer based, with the most widespread being polyamide thin film composite (PATFC) membranes.^{295,296} PATFC membranes are favoured due to their relatively wide operating pH (0–11), temperature (0–45 °C), and resilience to biological attack. The main problems for RO durability are its susceptibility to biofouling and chlorine oxidation.²⁹⁷ Blocked microbes and organisms accumulate on the saltwater membrane side, blocking the membrane pores and decreasing the active separation area. In order to prevent biofouling, the incoming water is often chlorinated to disinfect any microorganisms. However, addition of chlorine exacerbates membrane degradation by exposing it to oxidation attack. Commercial RO membranes are known to tolerate 1000 ppm h of Cl⁻, hence continuous exposure of chlorine at concentrations <0.1 ppm is acceptable.²⁹⁸ Most commercial RO membranes have a maximum operating temperature of 45–50 °C.²⁹⁵ The polyamide in PATFC, for instance, undergoes annealing above 45 °C causing a lower water permeation rate through the membrane over time.

5.2.2 Ion exchange resins. In order to achieve water quality >1 MΩ cm, an ion exchange resin is often added after the RO. An ion exchange resin works by exchanging various cations and anions present in the aqueous phase with other ions immobilised on solid resin beads, typically H⁺ and OH⁻ for high purity water. The beads consist of a matrix of functional polymers, such as polystyrene, crosslinked with divinylbenzene or an acrylic polymer. In general, there are 4 types of ion exchange resin: (i) Strong Acid Cation (SAC), (ii) Weak Acid Cation (WAC), (iii) Strong Base Anion (SBA), and (iv) Weak Base Anion (WBA). SAC and SBA are mostly used for complete demineralisation, which is the type of application for water electrolysis. The ion exchange site for SAC ion exchange resin is usually sulphonic acid (SO₃H) while for SBA ion exchange resin quaternary ammonium hydroxide groups of varying types are used. These are the same ion exchange groups that are used for membranes in PEMWES and AEMWES.

While fresh feed water is easily supplied at ambient temperature it is common in PEMWE systems to have purification in the recirculation loop fed from the outlet of the stack. A SAC resin can operate reliably above 100 °C. The bottleneck in ion exchange resin temperature stability is the reactivity of cationic headgroups in SBAs in the presence of hydroxide anions. As is very well known to researchers working with AEM materials, high temperatures facilitate degradation of these materials. Some commercial anion exchange resins can tolerate temperatures up to 70 °C but this may still be the most limiting component in PEMWE systems with respect to temperature, though we note that higher temperatures may compromise membrane mechanical integrity through creep deformation.⁷ Enhanced thermal stability may eventually be obtained by employing the techniques recently developed to improve the stability of AEM materials or potentially using non-hydroxide

anions that would not demineralise the water, but which would also not adversely impact performance.

5.3 Mitigation and recovery strategies

In the event of a contamination event or impurity accumulation, it may be worthwhile to attempt to recover the performance of an electrolyser. This section will suggest a few possible avenues to recover from impurity contamination in PEMWES, AWES, and AEMWES. To the best of our knowledge none of these are used in practice.

Contamination of the ion conducting phase of PEMWES, AWES and AEMWES is likely to be reversible. Metallic cation contamination of the PFSA in PEMWES may be removed by immersing in an acidic solution, such as 0.5–1.0 M H₂SO₄, to re-protonate the poisoned ionomer.¹²⁴ A novel approach to re-protonate the ionomer *in situ* is by flowing carbon dioxide at moderate pressures through the anode water tank to form carbonic acid.¹²³ Pressurising with carbon dioxide for 36 h was found to recover 95% cell voltage efficiency after poisoning by Fe²⁺ ions. In AWES and AEMWES using recirculating electrolyte an ion exchange resin cannot be placed inside the recirculation loop. Recovery from any contamination is therefore likely to require the replacement of the electrolyte. As with PEMWES, anionic contamination of AEM and ionomer may be reversible. Light carbonate contamination of the membrane or ionomer is expected to be self-reversing with operation at high current densities as OH⁻ is constantly generated at the cathode side²⁴¹ and would eventually purge carbonate and bicarbonate from the system as carbon dioxide. Other soluble anionic impurities may be flushed with repeated changes of electrolyte. If insoluble metal oxide precipitates are formed, then these may be removed by purging the system with an acidic solution before the electrolyte is replaced, though this may damage electrode materials.

At the catalyst layer, the chemistry of specific impurities will require different methods to clean the catalyst. Many ionic impurities be removed by oxidation or repeated rinsing of the system. Adsorbed halide anions on any catalyst surfaces may be desorbed by either oxidising at very positive potentials (>1 V *vs.* RHE) to oxidise the halide or by holding the catalyst at reducing potentials to evolve copious volumes of hydrogen. These techniques may require the cell potentials to be reversed, all techniques which involve potential excursions for the electrode may have a detrimental impact on catalyst longevity, but it may be relevant when contamination is critical. Similarly, high potentials may resolve heavy contamination from organic compounds *via* oxidation. Also noteworthy here are attempts to produce catalysts which are selective to OER in the presence of chlorine.²⁰⁹ Metallic or insoluble salt impurities may require removal by flushing the system with a solution which adjusts the pH into a range where such impurities are soluble (for example an acidic solution to remove iron precipitates). The challenge in all cases is to remove the impurity while avoiding, as far as possible, damage to the electrolyser components. This will require careful calibration of conditions for the recovery step.

Impurities added to hydrogen as a result of contamination may be mitigated by hydrogen purification steps, and very low



concentrations of trace impurities such as carbon dioxide and halides are tolerated in hydrogen fuel specifications, as discussed above. Radical formation may be mitigated using techniques applied in PEMFCs. The first method is to limit the crossover of hydrogen and oxygen by introducing a layer of recombination catalyst into the membrane or catalyst layer.^{133,299} Another method is to introduce a radical scavenger or peroxide decomposition catalyst inside the membrane to reduce the concentration of radicals that cause damage. Metal oxides such as titanium dioxide, zirconium dioxide, and cerium dioxide have proven their utility in suppressing the release of F⁻ under PEMFC operation.³⁰⁰

5.4 Summary and outstanding research challenges

5.4.1 Summary. The most direct method to study the impact of impurities is to introduce them during device testing at the single cell, or stack level. The drawback of this approach is that complementary techniques are needed to deconvolute which component of the cell or stack is degraded and elucidate any mechanism of operation. The applicable techniques highlighted here are summarised in Table 4.

In the liquid phase, the most commonly applied technique is ICP-MS, which can detect metallic elements down to the ppb level. However, most experiments use ICP-MS *ex situ*, meaning the water or electrolyte is collected at certain time intervals and analysis is performed afterwards. SFC ICP-MS offers an *operando* solution by connecting the cell directly with an ICP-MS. When the desired impurity is in ionic form, ion chromatography and polarography can work well; concentrations down to the ppb level can be measured. When the concentration is slightly higher, pH electrodes and ion sensitive electrodes can provide a cheaper and more robust alternative for *operando* measurement of impurity in the water.

Online gas phase analysis is rarely performed to study impurities; however, gas chromatography is the most common analytical technique applied, with different detectors and columns able to measure a range of gas phase impurities down to the ppb level. In the half cell setting, DEMS and EC-MS can be employed to track rapid evolution of side products semi-quantitatively. The RRDE, in this case, can also be used to monitor gaseous impurities that are generated as side-products at the anode.

Table 4 Summary of characterisation techniques for studying the impact of impurities on electrolyzers

Technique	Possible purpose	Detection limits
Rotating disc electrode	Impact of impurity on catalyst performance ^{260,261}	Not applicable
Single cell testing	Impact of impurity on CCM level performance ¹²⁴	Not applicable
Stack or system level testing	Impact of impurity on overall stack performance ⁴⁷	Not applicable
Rotating ring disc electrode	Side product measurement ^{147,182}	ppm (depending on molecule)
Electrochemical quartz crystal microbalance	Impact of change in mass due to impurity on electrode performance ⁸⁸	ng cm ⁻²
Attenuated total reflectance-Fourier transform infrared spectroscopy	Detecting change in functional group structure to study impact of impurities on membrane degradation ²⁰⁷	Not applicable
Ultraviolet-visible spectrophotometry	Concentration of impurities ²⁶⁷	ppb to ppm (depending on molecule)
X-ray absorption spectroscopy	Structure of catalyst surface ¹⁵¹	~5% by weight (depends on the element)
X-ray photoelectron spectroscopy	Surface characterization of catalysts ^{267,270,271}	0.1–1 at.%
Inductively coupled plasma-mass spectroscopy	Liquid phase quantification of metallic elements ^{68,278}	ppb
Scanning flow cell-inductively coupled mass spectrometry	<i>Operando</i> concentration detection ²⁷⁹	ppb
pH monitoring	Indication of impurity or side reaction taking place ¹²¹	Not applicable
Ion chromatography	Concentration measurement of ions ^{28,141,281}	ppb
Polarography	Concentration measurement of a specific ion (e.g. Pt ²⁺) ^{66,282}	ppb
Ion selective electrodes	Concentration measurement of a specific ion (e.g. F ⁻) ³⁰¹	ppb to ppm
Water conductivity	Conductivity measurement of feed water ^{210,287,288}	1 μS cm ⁻¹
Total organic carbon	Concentration of organic carbon ^{210,290}	5 mgC L ⁻¹
Gas chromatography	Concentration measurement of a molecule ^{196,291}	ppm
Differential electrochemical mass spectrometry	Qualitative concentration measurement of product and intermediate product ^{209,292}	Semi-quantitative
On chip electrochemical mass spectrometry	Quantitative concentration measurement of product and intermediate product ²⁹³	pmol s ⁻¹
Scanning electron microscopy-energy dispersive spectroscopy	Impurity distribution in CCM ⁶⁸	>0.1 wt%
Electron probe microanalysis	Impurity distribution in CCM ^{124,272}	ppm
Neutron/X-ray imaging	<i>Operando</i> distribution of impurity in the cell ^{273–276}	Semi-quantitative
Nanoscale secondary ion mass spectrometry	Impurity distribution in CCM ²⁷⁷	ppb to ppm



CCM characterisation is mostly performed *post mortem*, with the exception of online neutron imaging of impurities. Neutron imaging is a powerful *operando* tool that can show the distribution and poisoning behaviour of certain impurities within the CCM; however, this is usually limited to less relevant ions such as Gd^{3+} that have a sufficient neutron scattering cross-section. In *post mortem* testing the cell can be disassembled and individual components observed using SEM/EDX to map the distribution of impurities within the cell. EDX can only measure impurities at quite high concentrations, so for lower concentrations other techniques must be used. XPS and EPMA have been reported to be sensitive down to the ppm level while nano SIMS can go to ppb levels. Overall, the choice of technique depends on the nature and concentration of impurity present.

Water purification is a mature technology and there is little information available in the literature on the specific, proprietary systems implemented in commercial systems, but a combination of RO and ion exchange resin technologies is typical where high purity water is required.

The mitigation and recovery strategies presented are mostly theoretical processes that may be applied to recover electrolyzers in the event of severe poisoning or when the stack feed water is known to be impure. Processes exist for recovering poisoning by most types of impurity at low concentrations, but the processes may themselves cause damage to the electrolyser.

5.4.2 Outstanding research challenges. The study of the impact of impurities is relatively rare in the electrolyser field and the most commonly used approaches involve application of general-purpose analytical techniques, which have proven very powerful. Two broad challenges present themselves. The first is to measure the impact of low concentrations of various impurities on the lifetime of commercially relevant cells and systems, operating at relevant conditions. Most studies are necessarily limited to high-concentration impurity experiments to accelerate phenomena and make them easily observable in short duration experiments; however, in service electrolyzers are likely to be operated for 10000 s of hours and exposed to much lower concentrations of impurities. Without more long-term data and data at lower concentrations it will be challenging to understand the real significance of different impurities and their induced degradation mechanisms. The second challenge is to increase the time and spatial resolution of measurement techniques used to probe different mechanisms; this is especially important as evidence suggests that it is transient operation that results in the most significant cell degradation. For example, standard ICP-MS often has a low time resolution as it is an *ex situ* technique that requires manual sampling but the application of SFC ICP-MS, with a higher temporal resolution, effectively highlights the influence of potential transients.²⁰¹ Likewise, the application of neutron imaging was able to resolve the transient behaviour of ions *in situ*.¹²³

A clear challenge for water purification is to further reduce the capital and operating costs for purification. Ion exchange resins stable to increased temperature may be critical to support higher temperature operation of PEMWEs. Existing mitigation and recovery strategies are largely theoretical and need to be

demonstrated in real systems in order to properly evaluate their benefits, costs, and drawbacks.

6 Conclusions

As electrolyser developments continue and long-term, large-scale deployment ramps up, the limitations that impurities place on the economics, performance and durability of low temperature electrolyzers will become clearer. To date this has been a little studied area; the authors hope that this review will stimulate consideration of this area by summarising the state of the art, highlighting illustrative work in adjacent fields, and flagging outstanding research questions.

For PEMWEs, impurity testing at the cell level has only been performed explicitly for a handful of cations with others studied only in half-cell configurations or else their response inferred from results in analogous systems, such as PEMWEs. This is even more marked for anionic impurities, which have largely focused on Cl^- in half-cell tests; even here it is unclear how damaging chlorine evolution is when Cl^- is present at low concentrations for long time periods. The impact of simple organic molecules may be inferred from studies on alcohol-fed electrolyzers, but more complex molecules are likely to have an impact on durability.

The AWEs have been routinely operated for long periods suggesting that they are, in practice, robust to trace impurities. However, the limited studies on impurity tolerance mean that the impact on more modern higher-performing devices is unclear, as is the potentially unnecessary degree of conservatism in the water quality used. The use of lower catalyst loadings and more complex interfaces in AEMWEs is likely to make them less robust to impurities than AWEs, but again there is little hard information. Studies of the impact of impurities on AEMWEs are hindered by their ongoing development, which means that there is not yet a clear consensus on materials, design, or operation.

General questions relevant to all device types include methods for recovery from poisoning events, the stability/solubility of BoP materials in ultra-pure water, and the development of methods to assess the long-term stability towards impurities over short-time scales.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was funded by the National Measurement System of the UK's Department of Business, Energy & Industrial Strategy. JM acknowledges the Engineering and Physical Sciences Research Council for an Industrial CASE Studentship EP/S513635/1 with contributions from National Physical Laboratory and Johnson Matthey.



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